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Form PTO-1390 US DEPARTMENT OF COMMERCE PATENT AND IRADEMARK OF TRANSMITTAL LETTER TO THE UNITED STATES			ATTORNEY'S DOCKET NUMBER 2300-1481CIP				
DESIGNATED/ELEC CONCERNING A FIL		U.S. APPLICATION NO.					
INTERNATIONAL APPLICATION NO. INTERNATIONAL FI January 28,1999		ING DATE	PRIORITY DATE CLAIMED January 28, 1998				
TITLE OF INVENTION Novel Human Genes and Gene Expression Products II							
APPLICANT(S) FOR DO/EO/US Lewis T. Williams, et al.							
Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information: 1. X This is a FIRST submission of items concerning a filing under 35 U.S.C. 371.							
2 This is a SECOND or SUBSEQUE	ENT submission of items concerni	ing a filing under 35	U.S.C. 371.				
This express request to begin examination procedures (35 U.S.C. 371(f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and the PCT Articles 22 and 39(1).							
A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date.							
5. X A copy of the International Application as filed (35 U.S.C. 371(c)(2)): a is transmitted herewith (required only if not transmitted by the International Bureau). b has been transmitted by the International Bureau. c. X is not required, as the application was filed in the United States Receiving Office (RO/US).							
A translation of the International Application into English (35 U.S.C. 371(c)(2)).							
Amendments to the claims of the International Application under PCT Article 19 (37 U.S.C. 371(c)(3)): a are transmitted herewith (required only if not transmitted by the International Bureau). b have been transmitted by the International Bureau. c have not been made; however, the time limit for making such amendments has NOT expired. d X have not been made and will not be made.							
8 A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).							
9. X An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)). (UNSIGNED)							
10 A translation of the annexes to the	International Preliminary Examin	ation Report under P	CT Article 36 (35 U.S.C. 371(c)(5)).				
Items 11 to 16 below concern other document(s) or information included: 11 An Information Disclosure Statement under 37 CFR 1.97 and 1.98.							
2 An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.							
13 A FIRST preliminary amendment. A SECOND or SUBSEQUENT preliminary amendment.							
14 A substitute specification.	A substitute specification.						
15 A change of power of attorney and	A change of power of attorney and/or address letter.						
16 Other items or information:	Other items or information:						

U.S. APPLICATION NO (If know	rn, see 37 (FR 1 50)	INTERNATIONAL A PCT/US99/0161		A FIORNEY'S DOCKET NUMBER 2300-1481CIP			
17. X The following fee Basic National Fee (37)							
Neither international pr nor international search and international search	reliminary examination fee (and in fee (37 CFR 1.445(a)(2)) properties to the report not prepared by the	(37 CFR 1.482) baid to USPTO EPO or JPO	\$970.00				
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CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE				
Total Claims	21 - 20 =	01	X \$18.00	\$ 18.00			
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NOTE: Where an appropr must be filed and granted	riate time limit under 37 C	FR 1.494 or 1.495 has	not been met, a petition	to revive (37 CFR 1.13	37(a) or (b))		
BOZICEVIC, FIELD & FR. 285 Hamilton Avenue, Suite Palo Alto, California 94301 (650) 327-3400 Telephone (650) 327-3231 Facsimile	200		SIGNATURE Carol L. Francis NAME 36,513 REGISTRATION NUMI	A paul			

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NOVEL HUMAN GENES AND GENE EXPRESSION PRODUCTS II

Field of the Invention

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The present invention relates to novel polynucleotides, particularly to novel polynucleotides of human origin that are expressed in a selected cell type, are differentially expressed in one cell type relative to another cell type (e.g., in cancerous cells, or in cells of a specific tissue origin) and/or share homology to polynucleotides encoding a gene product having an identified functional domain and/or activity.

10 Background of the Invention

Identification of novel polynucleotides, particularly those that encode an expressed gene product, is important in the advancement of drug discovery, diagnostic technologies, and the understanding of the progression and nature of complex diseases such as cancer. Identification of genes expressed in different cell types isolated from sources that differ in disease state or stage, developmental stage, exposure to various environmental factors, the tissue of origin, the species from which the tissue was isolated, and the like is key to identifying the genetic factors that are responsible for the phenotypes associated with these various differences

This invention provides novel human polynucleotides, the polypeptides encoded by these polynucleotides, and the genes and proteins corresponding to these novel polynucleotides.

Summary of the Invention

This invention relates to novel human polynucleotides and variants thereof, their encoded polypeptides and variants thereof, to genes corresponding to these polynucleotides and to proteins expressed by the genes. The invention also relates to diagnostic and therapeutic agents employing such novel human polynucleotides, their corresponding genes or gene products, *e.g.*, these genes and proteins, including probes, antisense constructs, and antibodies. The polynucleotides of the invention correspond to a polynucleotide comprising the sequence information of at least one of SEQ ID NOS: 1-3544, 3546-4510, 4512-4725, 4727-4748, and 4750-5252, which for convenience sake is referred to herein as "SEQ ID NOS:1-5252."

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Accordingly, in one embodiment, the present invention features a library of polynucleotides, the library comprising the sequence information of at least one of "SEQ ID NOS:1-5252". In related aspects, the invention features a library provided on a nucleic acid array, or in a computer-readable format.

In one embodiment, the library is comprises a differentially expressed polynucleotide comprising a sequence selected from one of the differentially expressed polynucleotides disclosed herein. In specific related embodiments, the library comprises: 1) a polynucleotide that is differentially expressed in a human breast cancer cell, where the polynucleotide comprises a sequence selected from the group consisting of SEQ ID NOS:15, 36, 44, 45, 89, 146, 154, 159, 165, 172, 174, 183, 203, 261, 364, 366, 387, 419, 420, 496, 503, 510, 512, 529, 552, 560, 564, 570, 590, 606, 644, 646, 693, 707, 711, 726, 746, 754, 756, 875, 902, 921, 942, 990, 1095, 1104, 1122, 1131, 1142, 1170, 1184, 1205, 1286, 1289, 1354, 1387, 1435, 1535, 1751, 1764, 1777, 1795, 1860, 1869, 1882, 1890, 1915, 1933, 1934, 1979, 1980, 2007, 2023, 2040, 2059, 2223, 2245, 2300, 2325, 2409, 2462, 2488, 2486, and 2492; 2) a polynucleotide differentially expressed in a human colon cancer cell, where the polynucleotide comprises a sequence selected from the group consisting of SEQ ID NOS: , 33, 65, 228, 250, 252, 253, 280, 282, 355, 370, 387, 443, 460, 491, 545, 560, 581, 603, 680, 693, 703, 704, 716, 726, 746, 752, 753, 1095, 1104, 1205, 1241, 1264, 1354, 1387, 1401, 1442, 1514, 1734, 1742, 1780, 1851, 1899, 1915, 1954, 2024, 2066, 2262, and 2325; 3) a polynucleotide differentially expressed in a human lung cancer cell, where the polynucleotide comprises a sequence selected from the group consisting of SEQ ID NOS: 10, 54, 65, 171, 174, , 203, 252, 253, 254, , 285, 419, 420, 466, 491, 525, 526, 552, 571, 574, 590, 693, 700, 726, 742, 746, 861, 922, 990, 1088, 1288, 1355, 1417, 1422, 1444, 1454, 1570, 1597, 1979, 2007, 2024, 2034, 2038, 2126, and 2245; 4) a polynucleotide differentially expressed in growth factor-treated human microvascular endothelial cells (HMEC) relative to untreated HMEC, where the polynucleotide comprises a sequence selected from the group consisting of SEQ ID NOS:648, 1899, and 648; or 5) polynucleotides that are differentially expressed across multiple libraries, where the polynucleotide comprises a sequence selected from the group consisting of SEQ ID NOS: 65, 174, 203, 252, 253, 387, 419, 420, 491, 552, 560, 581, 590, 648, 693, 726, 746, 990, 1095, 1124, 1205, 1354, 1387, 1780, 1899, 1915, 1979, 2007, 2024, 2245, and 2325,

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In another aspect, the invention features an isolated polynucleotide comprising a nucleotide sequence having at least 90% sequence identity to an identifying sequence of "SEQ ID NOS:1-5252" or a degenerate variant thereof. In related aspects, the invention features recombinant host cells and vectors comprising the polynucleotides of the invention, as well as isolated polypeptides encoded by the polynucleotides of the invention and antibodies that specifically bind such polypeptides.

In one embodiment, the invention features an isolated polynucleotide comprising a sequence encoding a polypeptide of a protein family or having a functional domain selected from the group consisting of: 4 transmembrane segments integral membrane proteins, 7 transmembrane receptors (rhodopsin family or secretin family), eukaryotic aspartyl proteases, ATPases associated with various cellular activities (AAA), Bcl-2, cyclins, DEAD box protein family, DEAD/H helicase protein family, MAP kinase kinase protein family, novel 3'5'-cyclic nucleotide phosphodiesterases, protein kinases, ras protein family, G-protein alpha subunit, phorbol esters/diacylglycerol binding proteins, protein kinase, trypsin, protein tyrosine phosphatase, wnt family of developmental signaling proteins, WW/rsp5/WWP domain containing proteins, Ank repeat, basic region plus leucine zipper domain, bromodomain, eukaryotic thiol (cysteine) protease active site, EF-hand, ETS domain, type II fibronectin collagen binding domain, thioredoxin, homeobox domain, TNFR/NGFR family cysteine-rich region, WD domain/G-beta repeats, zinc finger (C2H2 type), zinc finger (CCHC class), and zinc finger (C3HC4 type). In a specific related embodiment, the invention features a polynucleotide comprising a sequence of one of the SEQ ID NOS: listed in Table 3 or Table 20.

In another aspect, the invention features a method of detecting differentially expressed genes correlated with a cancerous state of a mammalian cell, where the method comprises the step of detecting at least one differentially expressed gene product in a test sample derived from a cell suspected of being cancerous, where the gene product is encoded by a gene corresponding to a sequence of at least one of the differentially expressed polynucleotides disclosed herein. Detection of the differentially expressed gene product is correlated with a cancerous state of the cell from which the test sample was derived. In one embodiment, the detecting is by hybridization of the test sample to a

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reference array, wherein the reference array comprises an identifying sequence of at least one of the differentially expressed polynucleotides disclosed herein.

In one embodiment of the method of the invention, the cell is a breast tissue derived cell, and the differentially expressed gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS:15, 36, 44, 45, 89, 146, 154, 159, 165, 172, 174, 183, 203, 261, 364, 366, 387, 419, 420, 496, 503, 510, 512, 529, 552, 560, 564, 570, 590, 606, 644, 646, 693, 707, 711, 726, 746, 754, 756, 875, 902, 921, 942, 990, 1095, 1104, 1122, 1131, 1142, 1170, 1184, 1205, 1286, 1289, 1354, 1387, 1435, 1535, 1751, 1764, 1777, 1795, 1860, 1869, 1882, 1890, 1915, 1933, 1934, 1979, 1980, 2007, 2023, 2040, 2059, 2223, 2245, 2300, 2325, 2409, 2462, 2486 2488, and 2492.

In another embodiment of the method of the invention, the cell is a colon tissue derived cell, and differentially expressed gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS: 65, 228, 252, 253, 280, 355, 491, 581, 603, 680, 693, 716, 726, 746, 752, 753, 1241, 1264, 1401, 1442, 1514, 1851, 1915, 2024, 2066, 33, 250, 282, 370, 387, 443, 460, 545, 560, 703, 704, 1095, 1104, 1205, 1354, 1387, 1734, 1742, 1780, 1899, 1954, 2262, and 2325.

In yet another embodiment of the method of the invention, the cell is a lung tissue derived cell, and differentially expressed gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS:10, 54, 65, 171, 174, 203, 252, 253, 254, 285, 419, 420, 466, 491, 525, 526, 552, 571, 574, 590, 693, 700, 726, 742, 746, 861, 922, 990, 1088, 1288, 1355, 1417, 1422, 1444, 1454, 1570, 1597, 1979, 2007, 2024, 2034, 2038, 2126, and 2245.

In another embodiment, the cell is any of a lung, breast, or colon cell and the differentially expressed gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS:648 and 1899.

In still another embodiment, the cell is any of a breast, colon, or lung cell and the differentially expressed gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS: 65, 174, 203, 252, 253, 387, 419, 420, 491, 552, 560, 581, 590, 648, 693, 726, 746, 990, 1095, 1124, 1205, 1354, 1387, , 1780, 1899, 1915, 1979, 2007, 2024, 2245, and 2325.

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Other aspects and embodiments of the invention will be readily apparent to the ordinarily skilled artisan upon reading the description provided herein.

Detailed Description of the Invention

The invention relates to polynucleotides comprising the disclosed nucleotide sequences, to full length cDNA, mRNA and genes corresponding to these sequences, and to polypeptides and proteins encoded by these polynucleotides and genes.

Also included are polynucleotides that encode polypeptides and proteins encoded by the polynucleotides of the Sequence Listing. The various polynucleotides that can encode these polypeptides and proteins differ because of the degeneracy of the genetic code, in that most amino acids are encoded by more than one triplet codon. The identity of such codons is well-known in this art, and this information can be used for the construction of the polynucleotides within the scope of the invention.

Polynucleotides encoding polypeptides and proteins that are variants of the polypeptides and proteins encoded by the polynucleotides and related cDNA and genes are also within the scope of the invention. The variants differ from wild type protein in having one or more amino acid substitutions that either enhance, add, or diminish a biological activity of the wild type protein. Once the amino acid change is selected, a polynucleotide encoding that variant is constructed according to the invention.

The following detailed description describes the polynucleotide compositions encompassed by the invention, methods for obtaining cDNA or genomic DNA encoding a full-length gene product, expression of these polynucleotides and genes, identification of structural motifs of the polynucleotides and genes, identification of the function of a gene product encoded by a gene corresponding to a polynucleotide of the invention, use of the provided polynucleotides as probes and in mapping and in tissue profiling, use of the corresponding polypeptides and other gene products to raise antibodies, and use of the polynucleotides and their encoded gene products for therapeutic and diagnostic purposes.

I. <u>Polynucleotide Compositions</u>

The scope of the invention with respect to polynucleotide compositions includes, but is not necessarily limited to, polynucleotides having a sequence set forth in any one of

"SEQ ID NOS:1-5252"; polynucleotides obtained from the biological materials described herein or other biological sources (particularly human sources) by hybridization under stringent conditions (particularly conditions of high stringency); genes corresponding to the provided polynucleotides; variants of the provided polynucleotides and their corresponding genes, particularly those variants that retain a biological activity of the encoded gene product (e.g., a biological activity ascribed to a gene product corresponding to the provided polynucleotides as a result of the assignment of the gene product to a protein family(ies) and/or identification of a functional domain present in the gene product). Other nucleic acid compositions contemplated by and within the scope of the present invention will be readily apparent to one of ordinary skill in the art when provided with the disclosure here.

The invention features polynucleotides that are expressed in cells of human tissue, specifically human colon, breast, and/or lung tissue. Novel nucleic acid compositions of the invention of particular interest comprise a sequence set forth in any one of "SEQ ID NOS:1-5252" or an identifying sequence thereof. An "identifying sequence" is a contiguous sequence of residues at least about 10 nt to about 20 nt in length, usually at least about 50 nt to about 100 nt in length, that uniquely identifies a polynucleotide sequence, e.g., exhibits less than 90%, usually less than about 80% to about 85% sequence identity to any contiguous nucleotide sequence of more than about 20 nt. Thus, the subject novel nucleic acid compositions include full length cDNAs or mRNAs that encompass an identifying sequence of contiguous nucleotides from any one of "SEQ ID NOS:1-5252."

The polynucleotides of the invention also include polynucleotides having sequence similarity or sequence identity. Nucleic acids having sequence similarity are detected by hybridization under low stringency conditions, for example, at 50°C and 10XSSC (0.9 M saline/0.09 M sodium citrate) and remain bound when subjected to washing at 55°C in 1XSSC. Sequence identity can be determined by hybridization under stringent conditions, for example, at 50°C or higher and 0.1XSSC (9 mM saline/0.9 mM sodium citrate). Hybridization methods and conditions are well known in the art, see, e.g., U.S. Patent No. 5,707,829. Nucleic acids that are substantially identical to the provided polynucleotide sequences, e.g. allelic variants, genetically altered versions of the gene, etc., bind to the provided polynucleotide sequences ("SEQ ID NOS:1-5252") under stringent hybridization conditions. By using probes, particularly labeled probes of DNA sequences, one can

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isolate homologous or related genes. The source of homologous genes can be any species, e.g. primate species, particularly human; rodents, such as rats and mice; canines, felines, bovines, ovines, equines, yeast, nematodes, etc.

Preferably, hybridization is performed using at least 15 contiguous nucleotides of at least one of "SEQ ID NOS:1-5252." That is, when at least 15 contiguous nucleotides of one of the disclosed SEQ ID NOs. is used as a probe, the probe will preferentially hybridize with a gene or mRNA (of the biological material) comprising the complementary sequence, allowing the identification and retrieval of the nucleic acids of the biological material that uniquely hybridize to the selected probe. Probes from more than one SEQ ID NO. will hybridize with the same gene or mRNA if the cDNA from which they were derived corresponds to one mRNA. Probes of more than 15 nucleotides can be used, but 15 nucleotides represents enough sequence for unique identification.

The polynucleotides of the invention also include naturally occurring variants of the nucleotide sequences (e.g., degenerate variants, allelic variants, etc.). Variants of the polynucleotides of the invention are identified by hybridization of putative variants with nucleotide sequences disclosed herein, preferably by hybridization under stringent conditions. For example, by using appropriate wash conditions, variants of the polynucleotides of the invention can be identified where the allelic variant exhibits at most about 25-30% base pair mismatches relative to the selected polynucleotide probe. In general, allelic variants contain 15-25% base pair mismatches, and can contain as little as even 5-15%, or 2-5%, or 1-2% base pair mismatches, as well as a single base-pair mismatch.

The invention also encompasses homologs corresponding to the polynucleotides of "SEQ ID NOS:1-5252", where the source of homologous genes can be any mammalian species, e.g., primate species, particularly human; rodents, such as rats; canines, felines, bovines, ovines, equines, yeast, nematodes, etc. Between mammalian species, e.g., human and mouse, homologs have substantial sequence similarity, e.g., at least 75% sequence identity, usually at least 90%, more usually at least 95% between nucleotide sequences. Sequence similarity is calculated based on a reference sequence, which may be a subset of a larger sequence, such as a conserved motif, coding region, flanking region, etc. A reference sequence will usually be at least about 18 contiguous nt long, more usually at

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least about 30 nt long, and may extend to the complete sequence that is being compared. Algorithms for sequence analysis are known in the art, such as BLAST, described in Altschul *et al.*, *J. Mol. Biol.* (1990) 215:403-10.

In general, variants of the invention have a sequence identity greater than at least about 65%, preferably at least about 75%, more preferably at least about 85%, and can be greater than at least about 90% or more as determined by the Smith-Waterman homology search algorithm as implemented in MPSRCH program (Oxford Molecular). For the purposes of this invention, a preferred method of calculating percent identity is the Smith-Waterman algorithm, using the following. Global DNA sequence identity must be greater than 65% as determined by the Smith-Waterman homology search algorithm as implemented in MPSRCH program (Oxford Molecular) using an affine gap search with the following search parameters: gap open penalty, 12; and gap extension penalty, 1.

The subject nucleic acids can be cDNAs or genomic DNAs, as well as fragments thereof, particularly fragments that encode a biologically active gene product and/or are useful in the methods disclosed herein (e.g., in diagnosis, as a unique identifier of a differentially expressed gene of interest, etc.). The term "cDNA" as used herein is intended to include all nucleic acids that share the arrangement of sequence elements found in native mature mRNA species, where sequence elements are exons and 3' and 5' non-coding regions. Normally mRNA species have contiguous exons, with the intervening introns, when present, being removed by nuclear RNA splicing, to create a continuous open reading frame encoding a polypeptide of the invention.

A genomic sequence of interest comprises the nucleic acid present between the initiation codon and the stop codon, as defined in the listed sequences, including all of the introns that are normally present in a native chromosome. It can further include the 3' and 5' untranslated regions found in the mature mRNA. It can further include specific transcriptional and translational regulatory sequences, such as promoters, enhancers, etc., including about 1 kb, but possibly more, of flanking genomic DNA at either the 5' and 3' end of the transcribed region. The genomic DNA can be isolated as a fragment of 100 kbp or smaller; and substantially free of flanking chromosomal sequence. The genomic DNA flanking the coding region, either 3' and 5', or internal regulatory sequences as sometimes

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found in introns, contains sequences required for proper tissue, stage-specific, or diseasestate specific expression.

The nucleic acid compositions of the subject invention can encode all or a part of the subject polypeptides. Double or single stranded fragments can be obtained from the DNA sequence by chemically synthesizing oligonucleotides in accordance with conventional methods, by restriction enzyme digestion, by PCR amplification, *etc*. Isolated polynucleotides and polynucleotide fragments of the invention comprise at least about 10, about 15, about 20, about 35, about 50, about 100, about 150 to about 200, about 250 to about 300, or about 350 contiguous nucleotides selected from the polynucleotide sequences as shown in "SEQ ID NOS:1-5252." For the most part, fragments will be of at least 15 nt, usually at least 18 nt or 25 nt, and up to at least about 50 contiguous nt in length or more. In a preferred embodiment, the polynucleotide molecules comprise a contiguous sequence of at least twelve nucleotides selected from the group consisting of the polynucleotides shown in "SEQ ID NOS:1-5252."

Probes specific to the polynucleotides of the invention can be generated using the polynucleotide sequences disclosed in "SEQ ID NOS:1-5252." The probes are preferably at least about 12, 15, 16, 18, 20, 22, 24, or 25 nucleotide fragment of a corresponding contiguous sequence of "SEQ ID NOS:1-5252", and can be less than 2, 1, 0.5, 0.1, or 0.05 kb in length. The probes can be synthesized chemically or can be generated from longer polynucleotides using restriction enzymes. The probes can be labeled, for example, with a radioactive, biotinylated, or fluorescent tag. Preferably, probes are designed based upon an identifying sequence of a polynucleotide of one of "SEQ ID NOS:1-5252." More preferably, probes are designed based on a contiguous sequence of one of the subject polynucleotides that remain unmasked following application of a masking program for masking low complexity (e.g., XBLAST) to the sequence., i.e., one would select an unmasked region, as indicated by the polynucleotides outside the poly-n stretches of the masked sequence produced by the masking program.

The polynucleotides of the subject invention are isolated and obtained in substantial purity, generally as other than an intact chromosome. Usually, the polynucleotides, either as DNA or RNA, will be obtained substantially free of other naturally-occurring nucleic acid sequences, generally being at least about 50%, usually at least about 90% pure and are

typically "recombinant", e.g., flanked by one or more nucleotides with which it is not normally associated on a naturally occurring chromosome.

The polynucleotides of the invention can be provided as a linear molecule or within a circular molecule. They can be provided within autonomously replicating molecules (vectors) or within molecules without replication sequences. They can be regulated by their own or by other regulatory sequences, as is known in the art. The polynucleotides of the invention can be introduced into suitable host cells using a variety of techniques which are available in the art, such as transferrin polycation-mediated DNA transfer, transfection with naked or encapsulated nucleic acids, liposome-mediated DNA transfer, intracellular transportation of DNA-coated latex beads, protoplast fusion, viral infection, electroporation, gene gun, calcium phosphate-mediated transfection, and the like.

The subject nucleic acid compositions can be used to, for example, produce polypeptides, as probes for the detection of mRNA of the invention in biological samples (e.g., extracts of human cells) to generate additional copies of the polynucleotides, to generate ribozymes or antisense oligonucleotides, and as single stranded DNA probes or as triple-strand forming oligonucleotides. The probes described herein can be used to, for example, determine the presence or absence of the polynucleotide sequences as shown in "SEQ ID NOS:1-5252" or variants thereof in a sample. These and other uses are described in more detail below.

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<u>Use of Polynucleotides to Obtain Full-Length cDNA and Full-Length Human Gene and Promoter Region</u>

Full-length cDNA molecules comprising the disclosed polynucleotides are obtained as follows. A polynucleotide having a sequence of one of "SEQ ID NOS:1-5252", or a portion thereof comprising at least 12, 15, 18, or 20 nucleotides, is used as a hybridization probe to detect hybridizing members of a cDNA library using probe design methods, cloning methods, and clone selection techniques such as those described in U.S. Patent No. 5,654,173. Libraries of cDNA are made from selected tissues, such as normal or tumor tissue, or from tissues of a mammal treated with, for example, a pharmaceutical agent. Preferably, the tissue is the same as the tissue from which the polynucleotides of the invention were isolated, as both the polynucleotides described herein and the cDNA

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represent expressed genes. Most preferably, the cDNA library is made from the biological material described herein in the Examples. Alternatively, many cDNA libraries are available commercially. (Sambrook *et al.*, *Molecular Cloning: A Laboratory Manual, 2nd Ed.*, (1989) Cold Spring Harbor Press, Cold Spring Harbor, NY). The choice of cell type for library construction can be made after the identity of the protein encoded by the gene corresponding to the polynucleotide of the invention is known. This will indicate which tissue and cell types are likely to express the related gene, and thus represent a suitable source for the mRNA for generating the cDNA. Where the provided polynucleotides are isolated from cDNA libraries, the libraries are prepared from mRNA of human colon cells, more preferably, human colon cancer cells, even more preferably, from a highly metastatic colon cell, Km12L4-A.

Techniques for producing and probing nucleic acid sequence libraries are described, for example, in Sambrook *et al.*, *Molecular Cloning: A Laboratory Manual, 2nd Ed.*, (1989) Cold Spring Harbor Press, Cold Spring Harbor, NY. The cDNA can be prepared by using primers based on sequence from "SEQ ID NOS:1-5252." In one embodiment, the cDNA library can be made from only poly-adenylated mRNA. Thus, poly-T primers can be used to prepare cDNA from the mRNA.

Members of the library that are larger than the provided polynucleotides, and preferably that encompass the complete coding sequence of the native message, are obtained. In order to confirm that the entire cDNA has been obtained, RNA protection experiments are performed as follows. Hybridization of a full-length cDNA to an mRNA will protect the RNA from RNase degradation. If the cDNA is not full length, then the portions of the mRNA that are not hybridized will be subject to RNase degradation. This is assayed, as is known in the art, by changes in electrophoretic mobility on polyacrylamide gels, or by detection of released monoribonucleotides. Sambrook *et al.*, *Molecular Cloning: A Laboratory Manual, 2nd Ed.*, (1989) Cold Spring Harbor Press, Cold Spring Harbor, NY. In order to obtain additional sequences 5' to the end of a partial cDNA, 5' RACE (*PCR Protocols: A Guide to Methods and Applications*, (1990) Academic Press, Inc.) is performed.

Genomic DNA is isolated using the provided polynucleotides in a manner similar to the isolation of full-length cDNAs. Briefly, the provided polynucleotides, or portions

thereof, are used as probes to libraries of genomic DNA. Preferably, the library is obtained from the cell type that was used to generate the polynucleotides of the invention, but this is not essential. Most preferably, the genomic DNA is obtained from the biological material described herein in the Examples. Such libraries can be in vectors suitable for carrying large segments of a genome, such as P1 or YAC, as described in detail in Sambrook *et al.*, 9.4-9.30. In addition, genomic sequences can be isolated from human BAC libraries, which are commercially available from Research Genetics, Inc., Huntville, Alabama, USA, for example. In order to obtain additional 5' or 3' sequences, chromosome walking is performed, as described in Sambrook *et al.*, such that adjacent and overlapping fragments of genomic DNA are isolated. These are mapped and pieced together, as is known in the art, using restriction digestion enzymes and DNA ligase.

Using the polynucleotide sequences of the invention, corresponding full-length genes can be isolated using both classical and PCR methods to construct and probe cDNA libraries. Using either method, Northern blots, preferably, are performed on a number of cell types to determine which cell lines express the gene of interest at the highest level. Classical methods of constructing cDNA libraries are taught in Sambrook *et al.*, *supra*. With these methods, cDNA can be produced from mRNA and inserted into viral or expression vectors. Typically, libraries of mRNA comprising poly(A) tails can be produced with poly(T) primers. Similarly, cDNA libraries can be produced using the instant sequences as primers.

PCR methods are used to amplify the members of a cDNA library that comprise the desired insert. In this case, the desired insert will contain sequence from the full length cDNA that corresponds to the instant polynucleotides. Such PCR methods include gene trapping and RACE methods. Gene trapping entails inserting a member of a cDNA library into a vector. The vector then is denatured to produce single stranded molecules. Next, a substrate-bound probe, such a biotinylated oligo, is used to trap cDNA inserts of interest. Biotinylated probes can be linked to an avidin-bound solid substrate. PCR methods can be used to amplify the trapped cDNA. To trap sequences corresponding to the full length genes, the labeled probe sequence is based on the polynucleotide sequences of the invention. Random primers or primers specific to the library vector can be used to amplify the trapped cDNA. Such gene trapping techniques are described in Gruber *et al.*, WO

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95/04745 and Gruber *et al.*, U.S. Pat. No. 5,500,356. Kits are commercially available to perform gene trapping experiments from, for example, Life Technologies, Gaithersburg, Maryland, USA.

"Rapid amplification of cDNA ends," or RACE, is a PCR method of amplifying cDNAs from a number of different RNAs. The cDNAs are ligated to an oligonucleotide linker, and amplified by PCR using two primers. One primer is based on sequence from the instant polynucleotides, for which full length sequence is desired, and a second primer comprises sequence that hybridizes to the oligonucleotide linker to amplify the cDNA. A description of this methods is reported in WO 97/19110. In preferred embodiments of RACE, a common primer is designed to anneal to an arbitrary adaptor sequence ligated to cDNA ends (Apte and Siebert, *Biotechniques* (1993) 15:890-893; Edwards et al., Nuc. Acids Res. (1991) 19:5227-5232). When a single gene-specific RACE primer is paired with the common primer, preferential amplification of sequences between the single gene specific primer and the common primer occurs. Commercial cDNA pools modified for use in RACE are available.

Another PCR-based method generates full-length cDNA library with anchored ends without needing specific knowledge of the cDNA sequence. This method is described in WO 96/40998.

The promoter region of a gene generally is located 5' to the initiation site for RNA polymerase II. Hundreds of promoter regions contain the "TATA" box, a sequence such as TATTA or TATAA, which is sensitive to mutations. The promoter region can be obtained by performing 5' RACE using a primer from the coding region of the gene. Alternatively, the cDNA can be used as a probe for the genomic sequence, and the region 5' to the coding region is identified by "walking up." If the gene is highly expressed or differentially expressed, the promoter from the gene can be of use in a regulatory construct for a heterologous gene.

Once the full-length cDNA or gene is obtained, DNA encoding variants can be prepared by site-directed mutagenesis, described in detail in Sambrook *et al.*, 15.3-15.63. The choice of codon or nucleotide to be replaced can be based on disclosure herein on optional changes in amino acids to achieve altered protein structure and/or function.

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As an alternative method to obtaining DNA or RNA from a biological material, nucleic acid comprising nucleotides having the sequence of one or more polynucleotides of the invention can be synthesized. Thus, the invention encompasses nucleic acid molecules ranging in length from 15 nucleotides (corresponding to at least 15 contiguous nucleotides of one of "SEQ ID NOS:1-5252") up to a maximum length suitable for one or more biological manipulations, including replication and expression, of the nucleic acid molecule. The invention includes but is not limited to (a) nucleic acid having the size of a full gene, and comprising at least one of "SEQ ID NOS:1-5252;"; (b) the nucleic acid of (a) also comprising at least one additional gene, operably linked to permit expression of a fusion protein; (c) an expression vector comprising (a) or (b); (d) a plasmid comprising (a) or (b); and (e) a recombinant viral particle comprising (a) or (b). Once provided with the polynucleotides disclosed herein, construction or preparation of (a) - (e) are well within the skill in the art.

The sequence of a nucleic acid comprising at least 15 contiguous nucleotides of at least any one of "SEQ ID NOS:1-5252,", preferably the entire sequence of at least any one of "SEQ ID NOS:1-5252," is not limited and can be any sequence of A, T, G, and/or C (for DNA) and A, U, G, and/or C (for RNA) or modified bases thereof, including inosine and pseudouridine. The choice of sequence will depend on the desired function and can be dictated by coding regions desired, the intron-like regions desired, and the regulatory regions desired. Where the entire sequence of any one of "SEQ ID NOS:1-5252" is within the nucleic acid, the nucleic acid obtained is referred to herein as a polynucleotide comprising the sequence of any one of "SEQ ID NOS:1-5252."

II. Expression of Polypeptide Encoded by Full-Length cDNA or Full-Length Gene

The provided polynucleotide (e.g., a polynucleotide having a sequence of one of "SEQ ID NOS:1-5252"), the corresponding cDNA, or the full-length gene is used to express a partial or complete gene product. Constructs of polynucleotides having sequences of "SEQ ID NOS:1-5252" can be generated synthetically. Alternatively, single-step assembly of a gene and entire plasmid from large numbers of oligodeoxyribonucleotides is described by, e.g., Stemmer et al., Gene (Amsterdam) (1995) 164(1):49-53. In this method, assembly PCR (the synthesis of long DNA sequences from

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large numbers of oligodeoxyribonucleotides (oligos)) is described. The method is derived from DNA shuffling (Stemmer, *Nature* (1994) 370:389-391), and does not rely on DNA ligase, but instead relies on DNA polymerase to build increasingly longer DNA fragments during the assembly process.

Appropriate polynucleotide constructs are purified using standard recombinant DNA techniques as described in, for example, Sambrook *et al.*, *Molecular Cloning: A Laboratory Manual, 2nd Ed.*, (1989) Cold Spring Harbor Press, Cold Spring Harbor, NY, and under current regulations described in United States Dept. of HHS, National Institute of Health (NIH) Guidelines for Recombinant DNA Research. The gene product encoded by a polynucleotide of the invention is expressed in any expression system, including, for example, bacterial, yeast, insect, amphibian and mammalian systems. Suitable vectors and host cells are described in U.S. Patent No. 5,654,173.

Bacteria. Expression systems in bacteria include those described in Chang et al., Nature (1978) 275:615; Goeddel et al., Nature (1979) 281:544; Goeddel et al., Nucleic Acids Res. (1980) 8:4057; EP 0 036,776; U.S. Patent No. 4,551,433; DeBoer et al., Proc. Natl. Acad. Sci. (USA) (1983) 80:21-25; and Siebenlist et al., Cell (1980) 20:269.

Yeast. Expression systems in yeast include those described in Hinnen et al., Proc. Natl. Acad. Sci. (USA) (1978) 75:1929; Ito et al., J. Bacteriol. (1983) 153:163; Kurtz et al., Mol. Cell. Biol. (1986) 6:142; Kunze et al., J. Basic Microbiol. (1985) 25:141; Gleeson et al., J. Gen. Microbiol. (1986) 132:3459; Roggenkamp et al., Mol. Gen. Genet. (1986) 202:302; Das et al., J. Bacteriol. (1984) 158:1165; De Louvencourt et al., J. Bacteriol. (1983) 154:737; Van den Berg et al., Bio/Technology (1990) 8:135; Kunze et al., J. Basic Microbiol. (1985) 25:141; Cregg et al., Mol. Cell. Biol. (1985) 5:3376; U.S. Patent Nos. 4,837,148 and 4,929,555; Beach and Nurse, Nature (1981) 300:706; Davidow et al., Curr.

Genet. (1985) 10:380; Gaillardin et al., Curr. Genet. (1985) 10:49; Ballance et al.,
 Biochem. Biophys. Res. Commun. (1983) 112:284-289; Tilburn et al., Gene (1983)
 26:205-221; Yelton et al., Proc. Natl. Acad. Sci. (USA) (1984) 81:1470-1474; Kelly and
 Hynes, EMBO J. (1985) 4:475479; EP 0 244,234; and WO 91/00357.

Insect Cells. Expression of heterologous genes in insects is accomplished as described in U.S. Patent No. 4,745,051; Friesen *et al.*, "The Regulation of Baculovirus Gene Expression", in: *The Molecular Biology Of Baculoviruses* (1986) (W. Doerfler, ed.);

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EP 0 127,839; EP 0 155,476; and Vlak et al., J. Gen. Virol. (1988) 69:765-776; Miller et al., Ann. Rev. Microbiol. (1988) 42:177; Carbonell et al., Gene (1988) 73:409; Maeda et al., Nature (1985) 315:592-594; Lebacq-Verheyden et al., Mol. Cell. Biol. (1988) 8:3129; Smith et al., Proc. Natl. Acad. Sci. (USA) (1985) 82:8844; Miyajima et al., Gene (1987) 58:273; and Martin et al., DNA (1988) 7:99. Numerous baculoviral strains and variants and corresponding permissive insect host cells from hosts are described in Luckow et al., Bio/Technology (1988) 6:47-55, Miller et al., Generic Engineering (1986) 8:277-279, and Maeda et al., Nature (1985) 315:592-594.

Mammalian Cells. Mammalian expression is accomplished as described in Dijkema et al., EMBO J. (1985) 4:761, Gorman et al., Proc. Natl. Acad. Sci. (USA) (1982) 79:6777, Boshart et al., Cell (1985) 41:521 and U.S. Patent No. 4,399,216. Other features of mammalian expression are facilitated as described in Ham and Wallace, Meth. Enz. (1979) 58:44, Barnes and Sato, Anal. Biochem. (1980) 102:255, U.S. Patent Nos. 4,767,704, 4,657,866, 4,927,762, 4,560,655, WO 90/103430, WO 87/00195, and U.S. RE 30,985.

Polynucleotide molecules comprising a polynucleotide sequence provided herein propagated by placing the molecule in a vector. Viral and non-viral vectors are used, including plasmids. The choice of plasmid will depend on the type of cell in which propagation is desired and the purpose of propagation. Certain vectors are useful for amplifying and making large amounts of the desired DNA sequence. Other vectors are suitable for expression in cells in culture. Still other vectors are suitable for transfer and expression in cells in a whole animal or person. The choice of appropriate vector is well within the skill of the art. Many such vectors are available commercially. The partial or full-length polynucleotide is inserted into a vector typically by means of DNA ligase attachment to a cleaved restriction enzyme site in the vector. Alternatively, the desired nucleotide sequence can be inserted by homologous recombination in vivo. Typically this is accomplished by attaching regions of homology to the vector on the flanks of the desired nucleotide sequence. Regions of homology are added by ligation of oligonucleotides, or by polymerase chain reaction using primers comprising both the region of homology and a portion of the desired nucleotide sequence, for example.

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The polynucleotides set forth in "SEQ ID NOS:1-5252" or their corresponding full-length polynucleotides are linked to regulatory sequences as appropriate to obtain the desired expression properties. These can include promoters (attached either at the 5' end of the sense strand or at the 3' end of the antisense strand), enhancers, terminators, operators, repressors, and inducers. The promoters can be regulated or constitutive. In some situations it may be desirable to use conditionally active promoters, such as tissue-specific or developmental stage-specific promoters. These are linked to the desired nucleotide sequence using the techniques described above for linkage to vectors. Any techniques known in the art can be used.

When any of the above host cells, or other appropriate host cells or organisms, are used to replicate and/or express the polynucleotides or nucleic acids of the invention, the resulting replicated nucleic acid, RNA, expressed protein or polypeptide, is within the scope of the invention as a product of the host cell or organism. The product is recovered by any appropriate means known in the art.

Once the gene corresponding to a selected polynucleotide is identified, its expression can be regulated in the cell to which the gene is native. For example, an endogenous gene of a cell can be regulated by an exogenous regulatory sequence as disclosed in U.S. Patent No. 5,641,670.

III. Identification of Functional and Structural Motifs of Novel Genes

A. Screening Polynucleotide Sequences and Amino Acid Sequences Against
Publicly Available Databases

Translations of the nucleotide sequence of the provided polynucleotides, cDNAs or full genes can be aligned with individual known sequences. Similarity with individual sequences can be used to determine the activity of the polypeptides encoded by the polynucleotides of the invention. For example, sequences that show similarity with a chemokine sequence can exhibit chemokine activities. Also, sequences exhibiting similarity with more than one individual sequence can exhibit activities that are characteristic of either or both individual sequences.

The full length sequences and fragments of the polynucleotide sequences of the nearest neighbors can be used as probes and primers to identify and isolate the full length

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sequence corresponding to provided polynucleotides. The nearest neighbors can indicate a tissue or cell type to be used to construct a library for the full-length sequences corresponding to the provided polynucleotides..

Typically, a selected polynucleotide is translated in all six frames to determine the best alignment with the individual sequences. The sequences disclosed herein in the Sequence Listing are in a 5' to 3' orientation and translation in three frames can be sufficient (with a few specific exceptions as described in the Examples). These amino acid sequences are referred to, generally, as query sequences, which will be aligned with the individual sequences. Databases with individual sequences are described in "Computer Methods for Macromolecular Sequence Analysis" *Methods in Enzymology* (1996) 266, Doolittle, Academic Press, Inc., a division of Harcourt Brace & Co., San Diego, California, USA. Databases include Genbank, EMBL, and DNA Database of Japan (DDBJ).

Query and individual sequences can be aligned using the methods and computer programs described above, and include BLAST, available over the world wide web at http://ww.ncbi.nlm.nih.gov/BLAST/. Another alignment algorithm is Fasta, available in the Genetics Computing Group (GCG) package, Madison, Wisconsin, USA, a wholly owned subsidiary of Oxford Molecular Group, Inc. Other techniques for alignment are described in Doolittle, *supra*. Preferably, an alignment program that permits gaps in the sequence is utilized to align the sequences. The Smith-Waterman is one type of algorithm that permits gaps in sequence alignments. See *Meth. Mol. Biol.* (1997) 70: 173-187. Also, the GAP program using the Needleman and Wunsch alignment method can be utilized to align sequences. An alternative search strategy uses MPSRCH software, which runs on a MASPAR computer. MPSRCH uses a Smith-Waterman algorithm to score sequences on a massively parallel computer. This approach improves ability to identify sequences that are distantly related matches, and is especially tolerant of small gaps and nucleotide sequence errors. Amino acid sequences encoded by the provided polynucleotides can be used to search both protein and DNA databases.

Results of individual and query sequence alignments can be divided into three categories, high similarity, weak similarity, and no similarity. Individual alignment results ranging from high similarity to weak similarity provide a basis for determining polypeptide activity and/or structure. Parameters for categorizing individual results include: percentage

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of the alignment region length where the strongest alignment is found, percent sequence identity, and p value.

The percentage of the alignment region length is calculated by counting the number of residues of the individual sequence found in the region of strongest alignment, *e.g.*, contiguous region of the individual sequence that contains the greatest number of residues that are identical to the residues of the corresponding region of the aligned query sequence. This number is divided by the total residue length of the query sequence to calculate a percentage. For example, a query sequence of 20 amino acid residues might be aligned with a 20 amino acid region of an individual sequence. The individual sequence might be identical to amino acid residues 5, 9-15, and 17-19 of the query sequence. The region of strongest alignment is thus the region stretching from residue 9-19, an 11 amino acid stretch. The percentage of the alignment region length is: 11 (length of the region of strongest alignment) divided by (query sequence length) 20 or 55%.

Percent sequence identity is calculated by counting the number of amino acid matches between the query and individual sequence and dividing total number of matches by the number of residues of the individual sequences found in the region of strongest alignment. Thus, the percent identity in the example above would be 10 matches divided by 11 amino acids, or approximately, 90.9%

P value is the probability that the alignment was produced by chance. For a single alignment, the p value can be calculated according to Karlin et al., Proc. Natl. Acad. Sci. (1990) 87:2264 and Karlin et al., Proc. Natl. Acad. Sci. (1993) 90. The p value of multiple alignments using the same query sequence can be calculated using an heuristic approach described in Altschul et al., Nat. Genet. (1994) 6:119. Alignment programs such as BLAST program can calculate the p value.

Another factor to consider for determining identity or similarity is the location of the similarity or identity. Strong local alignment can indicate similarity even if the length of alignment is short. Sequence identity scattered throughout the length of the query sequence also can indicate a similarity between the query and profile sequences. The boundaries of the region where the sequences align can be determined according to Doolittle, *supra*; BLAST or FAST programs; or by determining the area where sequence identity is highest.

High Similarity. In general, in alignment results considered to be of high similarity, the percent of the alignment region length is typically at least about 55% of total length query sequence; more typically, at least about 58%; even more typically; at least about 60% of the total residue length of the query sequence. Usually, percent length of the alignment region can be as much as about 62%; more usually, as much as about 64%; even more usually, as much as about 66%. Further, for high similarity, the region of alignment, typically, exhibits at least about 75% of sequence identity; more typically, at least about 78%; even more typically; at least about 80% sequence identity. Usually, percent sequence identity can be as much as about 82%; more usually, as much as about 84%; even more usually, as much as about 86%.

The p value is used in conjunction with these methods. If high similarity is found, the query sequence is considered to have high similarity with a profile sequence when the p value is less than or equal to about 10^{-2} ; more usually; less than or equal to about 10^{-3} ; even more usually; less than or equal to about 10^{-4} . More typically, the p value is no more than about 10^{-5} ; more typically; no more than or equal to about 10^{-10} ; even more typically; no more than or equal to about 10^{-15} for the query sequence to be considered high similarity.

Weak Similarity. In general, where alignment results considered to be of weak similarity, there is no minimum percent length of the alignment region nor minimum length of alignment. A better showing of weak similarity is considered when the region of alignment is, typically, at least about 15 amino acid residues in length; more typically, at least about 20; even more typically; at least about 25 amino acid residues in length.

Usually, length of the alignment region can be as much as about 30 amino acid residues; more usually, as much as about 40; even more usually, as much as about 60 amino acid residues. Further, for weak similarity, the region of alignment, typically, exhibits at least about 35% of sequence identity; more typically, at least about 40%; even more typically; at least about 45% sequence identity. Usually, percent sequence identity can be as much as about 50%; more usually, as much as about 55%; even more usually, as much as about 60%.

If low similarity is found, the query sequence is considered to have weak similarity with a profile sequence when the p value is usually less than or equal to about 10⁻²; more usually; less than or equal to about 10⁻³; even more usually; less than or equal to about 10⁻⁴. More

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typically, the p value is no more than about 10⁻⁵; more usually; no more than or equal to about 10⁻¹⁰; even more usually; no more than or equal to about 10⁻¹⁵ for the query sequence to be considered weak similarity.

Similarity Determined by Sequence Identity Alone. Sequence identity alone can be used to determine similarity of a query sequence to an individual sequence and can indicate the activity of the sequence. Such an alignment, preferably, permits gaps to align sequences. Typically, the query sequence is related to the profile sequence if the sequence identity over the entire query sequence is at least about 15%; more typically, at least about 20%; even more typically, at least about 25%; even more typically, at least about 50%. Sequence identity alone as a measure of similarity is most useful when the query sequence is usually, at least 80 residues in length; more usually, 90 residues; even more usually, at least 95 amino acid residues in length. More typically, similarity can be concluded based on sequence identity alone when the query sequence is preferably 100 residues in length; more preferably, 120 residues in length; even more preferably, 150 amino acid residues in length.

<u>Determining Activity from Alignments with Profile and Multiple Aligned</u>
<u>Sequences.</u> Translations of the provided polynucleotides can be aligned with amino acid profiles that define either protein families or common motifs. Also, translations of the provided polynucleotides can be aligned to multiple sequence alignments (MSA) comprising the polypeptide sequences of members of protein families or motifs. Similarity or identity with profile sequences or MSAs can be used to determine the activity of the gene products (*e.g.*, polypeptides) encoded by the provided polynucleotides or corresponding cDNA or genes. For example, sequences that show an identity or similarity with a chemokine profile or MSA can exhibit chemokine activities.

Profiles can designed manually by (1) creating an MSA, which is an alignment of the amino acid sequence of members that belong to the family and (2) constructing a statistical representation of the alignment. Such methods are described, for example, in Birney et al., Nucl. Acid Res. (1996) 24(14): 2730-2739. MSAs of some protein families and motifs are publicly available. For example, http://genome.wustl.edu/Pfam/ includes MSAs of 547 different families and motifs. These MSAs are described also in Sonnhammer et al., Proteins (1997) 28: 405-420. Other sources over the world wide web

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include the site at http://www.embl-heidelberg.de/argos/ali/ali.htm1; alternatively, a message can be sent to ALI@EMBL-HEIDELBERG.DE for the information. A brief description of these MSAs is reported in Pascarella et al., Prot. Eng. (1996) 9(3):249-251. Techniques for building profiles from MSAs are described in Sonnhammer et al., supra; Birney et al., supra; and "Computer Methods for Macromolecular Sequence Analysis," Methods in Enzymology (1996) 266, Doolittle, Academic Press, Inc., a division of Harcourt Brace & Co., San Diego, California, USA.

Similarity between a query sequence and a protein family or motif can be determined by (a) comparing the query sequence against the profile and/or (b) aligning the query sequence with the members of the family or motif. Typically, a program such as Searchwise is used to compare the query sequence to the statistical representation of the multiple alignment, also known as a profile. The program is described in Birney *et al.*, *supra*. Other techniques to compare the sequence and profile are described in Sonnhammer *et al.*, *supra* and Doolittle, *supra*.

Next, methods described by Feng et al., J. Mol. Evol. (1987) 25:351 and Higgins et al., CABIOS (1989) 5:151 can be used align the query sequence with the members of a family or motif, also known as a MSA. Computer programs, such as PILEUP, can be used. See Feng et al., infra. In general, the following factors are used to determine if a similarity between a query sequence and a profile or MSA exists: (1) number of conserved residues found in the query sequence, (2) percentage of conserved residues found in the query sequence, (3) number of frameshifts, and (4) spacing between conserved residues.

Some alignment programs that both translate and align sequences can make any number of frameshifts when translating the nucleotide sequence to produce the best alignment. The fewer frameshifts needed to produce an alignment, the stronger the similarity or identity between the query and profile or MSAs. For example, a weak similarity resulting from no frameshifts can be a better indication of activity or structure of a query sequence, than a strong similarity resulting from two frameshifts. Preferably, three or fewer frameshifts are found in an alignment; more preferably two or fewer frameshifts; even more preferably, no frameshifts are found in an alignment of query and profile or MSAs.

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Conserved residues are those amino acids found at a particular position in all or some of the family or motif members. For example, most chemokines contain four conserved cysteines. Alternatively, a position is considered conserved if only a certain class of amino acids is found in a particular position in all or some of the family members. For example, the N-terminal position can contain a positively charged amino acid, such as lysine, arginine, or histidine.

Typically, a residue of a polypeptide is conserved when a class of amino acids or a single amino acid is found at a particular position in at least about 40% of all class members; more typically, at least about 50%; even more typically, at least about 60% of the members. Usually, a residue is conserved when a class or single amino acid is found in at least about 70% of the members of a family or motif; more usually, at least about 80%; even more usually, at least about 95%.

A residue is considered conserved when three unrelated amino acids are found at a particular position in the some or all of the members; more usually, two unrelated amino acids. These residues are conserved when the unrelated amino acids are found at particular positions in at least about 40% of all class member; more typically, at least about 50%; even more typically, at least about 60% of the members. Usually, a residue is conserved when a class or single amino acid is found in at least about 70% of the members of a family or motif; more usually, at least about 80%; even more usually, at least about 90%; even more usually, at least about 95%.

A query sequence has similarity to a profile or MSA when the query sequence comprises at least about 25% of the conserved residues of the profile or MSA; more usually, at least about 30%; even more usually; at least about 40%. Typically, the query sequence has a stronger similarity to a profile sequence or MSA when the query sequence comprises at least about 45% of the conserved residues of the profile or MSA; more typically, at least about 50%; even more typically; at least about 55%.

B. <u>Screening Polynucleotide and Amino Acid Sequences Against Protein</u>

Profiles

The identify and function of the gene that correlates to a polynucleotide described herein can be determined by screening the polynucleotides or their corresponding amino acid sequences against profiles of protein families. Such profiles focus on common

structural motifs among proteins of each family. Publicly available profiles are described above in Section IVA. Additional or alternative profiles are described below.

In comparing a novel polynucleotide with known sequences, several alignment tools are available. Examples include PileUp, which creates a multiple sequence alignment, and is described in Feng et al., J. Mol. Evol. (1987) 25:351. Another method, GAP, uses the alignment method of Needleman et al., J. Mol. Biol. (1970) 48:443. GAP is best suited for global alignment of sequences. A third method, BestFit, functions by inserting gaps to maximize the number of matches using the local homology algorithm of Smith et al., Adv. Appl. Math. (1981) 2:482.

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C. <u>Identification of Secreted & Membrane-Bound Polypeptides</u>

Both secreted and membrane-bound polypeptides of the present invention are of particular interest. For example, levels of secreted polypeptides can be assayed in body fluids that are convenient, such as blood, urine, prostatic fluid and semen. Membrane-bound polypeptides are useful for constructing vaccine antigens or inducing an immune response. Such antigens would comprise all or part of the extracellular region of the membrane-bound polypeptides. Because both secreted and membrane-bound polypeptides comprise a fragment of contiguous hydrophobic amino acids, hydrophobicity predicting algorithms can be used to identify such polypeptides.

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A signal sequence is usually encoded by both secreted and membrane-bound polypeptide genes to direct a polypeptide to the surface of the cell. The signal sequence usually comprises a stretch of hydrophobic residues. Such signal sequences can fold into helical structures. Membrane-bound polypeptides typically comprise at least one transmembrane region that possesses a stretch of hydrophobic amino acids that can transverse the membrane. Some transmembrane regions also exhibit a helical structure. Hydrophobic fragments within a polypeptide can be identified by using computer algorithms. Such algorithms include Hopp & Woods, *Proc. Natl. Acad. Sci. USA* (1981) 78:3824-3828; Kyte & Doolittle, *J. Mol. Biol.* (1982) 157: 105-132; and RAOAR algorithm, Degli Esposti *et al.*, *Eur. J. Biochem.* (1990) 190: 207-219.

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Another method of identifying secreted and membrane-bound polypeptides is to translate the polynucleotides of the invention in all six frames and determine if at least 8

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contiguous hydrophobic amino acids are present. Those translated polypeptides with at least 8; more typically, 10; even more typically, 12 contiguous hydrophobic amino acids are considered to be either a putative secreted or membrane bound polypeptide.

Hydrophobic amino acids include alanine, glycine, histidine, isoleucine, leucine, lysine, methionine, phenylalanine, proline, threonine, tryptophan, tyrosine, and valine.

IV. <u>Identification of the Function of an Expression Product of a Full-Length Gene</u> Corresponding to a Polynucleotide

Ribozymes, antisense constructs, and dominant negative mutants can be used to determine function of the expression product of a gene corresponding to a polynucleotide provided herein. These methods and compositions are particularly useful where the provided novel polynucleotide exhibits no significant or substantial homology to a sequence encoding a gene of known function. Antisense molecules and ribozymes can be constructed from synthetic polynucleotides. Typically, the phosphoramidite method of oligonucleotide synthesis is used. See Beaucage et al., Tet. Lett. (1981) 22:1859 and U.S. Patent No. 4,668,777. Automated devices for synthesis are available to create oligonucleotides using this chemistry. Examples of such devices include Biosearch 8600, Models 392 and 394 by Applied Biosystems, a division of Perkin-Elmer Corp., Foster City, California, USA; and Expedite by Perceptive Biosystems, Framingham, Massachusetts, USA. Synthetic RNA, phosphate analog oligonucleotides, and chemically derivatized oligonucleotides can also be produced, and can be covalently attached to other molecules. RNA oligonucleotides can be synthesized, for example, using RNA phosphoramidites. This method can be performed on an automated synthesizer, such as Applied Biosystems, Models 392 and 394, Foster City, California, USA. See Applied Biosystems User Bulletin 53 and Ogilvie et al., Pure & Applied Chem. (1987) 59:325.

Phosphorothioate oligonucleotides can also be synthesized for antisense construction. A sulfurizing reagent, such as tetraethylthiruam disulfide (TETD) in acetonitrile can be used to convert the internucleotide cyanoethyl phosphite to the phosphorothioate triester within 15 minutes at room temperature. TETD replaces the iodine reagent, while all other reagents used for standard phosphoramidite chemistry

remain the same. Such a synthesis method can be automated using Models 392 and 394 by Applied Biosystems, for example.

Oligonucleotides of up to 200 nucleotides can be synthesized, more typically, 100 nucleotides, more typically 50 nucleotides; even more typically 30 to 40 nucleotides.

These synthetic fragments can be annealed and ligated together to construct larger

fragments. See, for example, Sambrook et al., supra.

A. Ribozymes

Trans-cleaving catalytic RNAs (ribozymes) are RNA molecules possessing endoribonuclease activity. Ribozymes are specifically designed for a particular target, and the target message must contain a specific nucleotide sequence. They are engineered to cleave any RNA species site-specifically in the background of cellular RNA. The cleavage event renders the mRNA unstable and prevents protein expression. Importantly, ribozymes can be used to inhibit expression of a gene of unknown function for the purpose of determining its function in an in vitro or in vivo context, by detecting the phenotypic effect.

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One commonly used ribozyme motif is the hammerhead, for which the substrate sequence requirements are minimal. Design of the hammerhead ribozyme is disclosed in Usman et al., Current Opin. Struct. Biol. (1996) 6:527. Ribozymes can also be prepared and used as described in Long et al., FASEB J. (1993) 7:25; Symons, Ann. Rev. Biochem. (1992) 61:641; Perrotta et al., Biochem. (1992) 31:16; Ojwang et al., Proc. Natl. Acad. Sci. (USA) (1992) 89:10802; and U.S. Patent No. 5,254,678. Ribozyme cleavage of HIV-I RNA is described in U.S. Patent No. 5,144,019; methods of cleaving RNA using ribozymes is described in U.S. Patent No. 5,116,742; and methods for increasing the specificity of ribozymes are described in U.S. Patent No. 5,225,337 and Koizumi et al., Nucleic Acid Res. (1989) 17:7059. Preparation and use of ribozyme fragments in a hammerhead structure are also described by Koizumi et al., Nucleic Acids Res. (1989) 17:7059. Preparation and use of ribozyme fragments in a hairpin structure are described by Chowrira and Burke, Nucleic Acids Res. (1992) 20:2835. Ribozymes can also be made by rolling transcription as described in Daubendiek and Kool, Nat. Biotechnol. (1997)

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The hybridizing region of the ribozyme can be modified or can be prepared as a branched structure as described in Horn and Urdea, *Nucleic Acids Res.* (1989) 17:6959. The basic structure of the ribozymes can also be chemically altered in ways familiar to those skilled in the art, and chemically synthesized ribozymes can be administered as synthetic oligonucleotide derivatives modified by monomeric units. In a therapeutic context, liposome mediated delivery of ribozymes improves cellular uptake, as described in Birikh *et al.*, *Eur. J. Biochem.* (1997) 245:1.

Using the polynucleotide sequences of the invention and methods known in the art, ribozymes are designed to specifically bind and cut the corresponding mRNA species. Ribozymes thus provide a means to inhibit the expression of any of the proteins encoded by the disclosed polynucleotides or their full-length genes. The full-length gene need not be known in order to design and use specific inhibitory ribozymes. In the case of a polynucleotide or full-length cDNA of unknown function, ribozymes corresponding to that nucleotide sequence can be tested in vitro for efficacy in cleaving the target transcript. Those ribozymes that effect cleavage in vitro are further tested in vivo. The ribozyme can also be used to generate an animal model for a disease, as described in Birikh *et al.*, *supra*. An effective ribozyme is used to determine the function of the gene of interest by blocking its transcription and detecting a change in the cell. Where the gene is found to be a mediator in a disease, an effective ribozyme is designed and delivered in a gene therapy for blocking transcription and expression of the gene.

Therapeutic and functional genomic applications of ribozymes proceed beginning with knowledge of a portion of the coding sequence of the gene to be inhibited. Thus, for many genes, a partial polynucleotide sequence provides adequate sequence for constructing an effective ribozyme. A target cleavage site is selected in the target sequence, and a ribozyme is constructed based on the 5' and 3' nucleotide sequences that flank the cleavage site. Retroviral vectors are engineered to express monomeric and multimeric hammerhead ribozymes targeting the mRNA of the target coding sequence. These monomeric and multimeric ribozymes are tested in vitro for an ability to cleave the target mRNA. A cell line is stably transduced with the retroviral vectors expressing the ribozymes, and the transduction is confirmed by Northern blot analysis and reverse-transcription polymerase chain reaction (RT-PCR). The cells are screened for inactivation of the target mRNA by

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such indicators as reduction of expression of disease markers or reduction of the gene product of the target mRNA.

B. Antisense

Antisense nucleic acids are designed to specifically bind to RNA, resulting in the formation of RNA-DNA or RNA-RNA hybrids, with an arrest of DNA replication, reverse transcription or messenger RNA translation. Antisense polynucleotides based on a selected polynucleotide sequence can interfere with expression of the corresponding gene. Antisense polynucleotides are typically generated within the cell by expression from antisense constructs that contain the antisense strand as the transcribed strand. Antisense polynucleotides based on the disclosed polynucleotides will bind and/or interfere with the translation of mRNA comprising a sequence complementary to the antisense polynucleotide. The expression products of control cells and cells treated with the antisense construct are compared to detect the protein product of the gene corresponding to the polynucleotide upon which the antisense construct is based. The protein is isolated and identified using routine biochemical methods.

Given the extensive background literature and clinical experience in antisense therapy, one skilled in the art can use selected polynucleotides of the invention as additional potential therapeutics. The choice of polynucleotide can be narrowed by first testing them for binding to "hot spot" regions of the genome of cancerous cells. If a polynucleotide is identified as binding to a "hot spot", testing the polynucleotide as an antisense compound in the corresponding cancer cells clearly is warranted.

C. Dominant Negative Mutations

As an alternative method for identifying function of the gene corresponding to a polynucleotide disclosed herein, dominant negative mutations are readily generated for corresponding proteins that are active as homomultimers. A mutant polypeptide will interact with wild-type polypeptides (made from the other allele) and form a non-functional multimer. Thus, a mutation is in a substrate-binding domain, a catalytic domain, or a cellular localization domain. Preferably, the mutant polypeptide will be overproduced. Point mutations are made that have such an effect. In addition, fusion of different polypeptides of various lengths to the terminus of a protein can yield dominant negative mutants. General strategies are available for making dominant negative mutants (see, e.g.,

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Herskowitz, *Nature* (1987) 329:219). Such techniques can be used to create loss of function mutations, which are useful for determining protein function.

V. Construction of Polypeptides of the Invention and Variants Thereof

The polypeptides of the invention include those encoded by the disclosed polynucleotides. These polypeptides can also be encoded by nucleic acids that, by virtue of the degeneracy of the genetic code, are not identical in sequence to the disclosed polynucleotides. Thus, the invention includes within its scope a polypeptide encoded by a polynucleotide having the sequence of any one of "SEQ ID NOS:1-5252" or a variant thereof.

In general, the term "polypeptide" as used herein refers to both the full length polypeptide encoded by the recited polynucleotide, the polypeptide encoded by the gene represented by the recited polynucleotide, as well as portions or fragments thereof. "Polypeptides" also includes variants of the naturally occurring proteins, where such variants are homologous or substantially similar to the naturally occurring protein, and can be of an origin of the same or different species as the naturally occurring protein (e.g., human, murine, or some other species that naturally expresses the recited polypeptide, usually a mammalian species). In general, variant polypeptides have a sequence that has at least about 80%, usually at least about 90%, and more usually at least about 98% sequence identity with a differentially expressed polypeptide of the invention, as measured by BLAST using the parameters described above. The variant polypeptides can be naturally or non-naturally glycosylated, i.e., the polypeptide has a glycosylation pattern that differs from the glycosylation pattern found in the corresponding naturally occurring protein.

The invention also encompasses homologs of the disclosed polypeptides (or fragments thereof) where the homologs are isolated from other species, *i.e.* other animal or plant species, where such homologs, usually mammalian species, *e.g.* rodents, such as mice, rats; domestic animals, *e.g.*, horse, cow, dog, cat; and humans. By homolog is meant a polypeptide having at least about 35%, usually at least about 40% and more usually at least about 60% amino acid sequence identity a particular differentially expressed protein as identified above, where sequence identity is determined using the BLAST algorithm, with the parameters described *supra*.

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In general, the polypeptides of the subject invention are provided in a non-naturally occurring environment, e.g. are separated from their naturally occurring environment. In certain embodiments, the subject protein is present in a composition that is enriched for the protein as compared to a control. As such, purified polypeptide is provided, where by purified is meant that the protein is present in a composition that is substantially free of non-differentially expressed polypeptides, where by substantially free is meant that less than 90%, usually less than 60% and more usually less than 50% of the composition is made up of non-differentially expressed polypeptides.

Also within the scope of the invention are variants; variants of polypeptides include mutants, fragments, and fusions. Mutants can include amino acid substitutions, additions or deletions. The amino acid substitutions can be conservative amino acid substitutions or substitutions to eliminate non-essential amino acids, such as to alter a glycosylation site, a phosphorylation site or an acetylation site, or to minimize misfolding by substitution or deletion of one or more cysteine residues that are not necessary for function. Conservative amino acid substitutions are those that preserve the general charge, hydrophobicity/hydrophilicity, and/or steric bulk of the amino acid substituted. For example, substitutions between the following groups are conservative: Gly/Ala, Val/Ile/Leu, Asp/Glu, Lys/Arg, Asn/Gln, Ser/Cys, Thr, and Phe/Trp/Tyr.

Variants can be designed so as to retain biological activity of a particular region of the protein (e.g., a functional domain and/or, where the polypeptide is a member of a protein family, a region associated with a consensus sequence). In a non-limiting example, Osawa et al., Biochem. Mol. Int. (1994) 34:1003, discusses the actin binding region of a protein from several different species. The actin binding regions of the these species are considered homologous based on the fact that they have amino acids that fall within "homologous residue groups." Homologous residues are judged according to the following groups (using single letter amino acid designations): STAG; ILVMF; HRK; DEQN; and FYW. For example, and S, a T, an A or a G can be in a position and the function (in this case actin binding) is retained.

Additional guidance on amino acid substitution is available from studies of protein evolution. Go et al, Int. J. Peptide Protein Res. (1980) 15:211, classified amino acid residue sites as interior or exterior depending on their accessibility. More frequent

substitution on exterior sites was confirmed to be general in eight sets of homologous protein families regardless of their biological functions and the presence or absence of a prosthetic group. Virtually all types of amino acid residues had higher mutabilities on the exterior than in the interior. No correlation between mutability and polarity was observed of amino acid residues in the interior and exterior, respectively. Amino acid residues were classified into one of three groups depending on their polarity: polar (Arg, Lys, His, Gln, Asn, Asp, and Glu); weak polar (Ala, Pro, Gly, Thr, and Ser), and nonpolar (Cys, Val, Met, Ile, Leu, Phe, Tyr, and Trp). Amino acid replacements during protein evolution were very conservative: 88% and 76% of them in the interior or exterior, respectively, were within the same group of the three. Inter-group replacements are such that weak polar residues are replaced more often by nonpolar residues in the interior and more often by polar residues on the exterior.

Additional guidance for production of polypeptide variants is provided in Querol et al., Prot. Eng. (1996) 9:265, which provides general rules for amino acid substitutions to enhance protein thermostability. New glycosylation sites can be introduced as discussed in Olsen and Thomsen, J. Gen. Microbiol. (1991) 137:579. An additional disulfide bridge can be introduced, as discussed by Perry and Wetzel, Science (1984) 226:555; Pantoliano et al., Biochemistry (1987) 26:2077; Matsumura et al., Nature (1989) 342:291; Nishikawa et al., Protein Eng. (1990) 3:443; Takagi et al., J. Biol. Chem. (1990) 265:6874; Clarke et al., Biochemistry (1993) 32:4322; and Wakarchuk et al., Protein Eng. (1994) 7:1379. Metal binding sites can be introduced, according to Toma et al., Biochemistry (1991) 30:97, and Haezerbrouck et al., Protein Eng. (1993) 6:643. Substitutions with prolines in loops can be made according to Masul et al., Appl. Env. Microbiol. (1994) 60:3579; and Hardy et al., FEBS Lett. 317:89.

Cysteine-depleted muteins are considered variants within the scope of the invention. These variants can be constructed according to methods disclosed in U.S. Patent No. 4,959,314, which discloses substitution of cysteines with other amino acids, and methods for assaying biological activity and effect of the substitution. Such methods are suitable for proteins according to this invention that have cysteine residues suitable for such substitutions, for example to eliminate disulfide bond formation.

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Variants also include fragments of the polypeptides disclosed herein, particularly biologically active fragments and/or fragments corresponding to functional domains. Fragments of interest will typically be at least about 10 aa to at least about 15 aa in length, usually at least about 50 aa in length, and can be as long as 300 aa in length or longer, but will usually not exceed about 1000 aa in length, where the fragment will have a stretch of amino acids that is identical to a polypeptide encoded by a polynucleotide having a sequence of any "SEQ ID NOS:1-5252", or a homolog thereof.

The protein variants described herein are encoded by polynucleotides that are within the scope of the invention. The genetic code can be used to select the appropriate codons to construct the corresponding variants.

VI. Computer-Related Embodiments

In general, a library of polynucleotides is a collection of sequence information, which information is provided in either biochemical form (e.g., as a collection of polynucleotide molecules), or in electronic form (e.g., as a collection of polynucleotide sequences stored in a computer-readable form, as in a computer system and/or as part of a computer program). The sequence information of the polynucleotides can be used in a variety of ways, e.g., as a resource for gene discovery, as a representation of sequences expressed in a selected cell type (e.g., cell type markers), and/or as markers of a given disease or disease state. In general, a disease marker is a representation of a gene product that is present in all cells affected by disease either at an increased or decreased level relative to a normal cell (e.g., a cell of the same or similar type that is not substantially affected by disease). For example, a polynucleotide sequence in a library can be a polynucleotide that represents an mRNA, polypeptide, or other gene product encoded by the polynucleotide, that is either overexpressed or underexpressed in a breast ductal cell affected by cancer relative to a normal (i.e., substantially disease-free) breast cell.

The nucleotide sequence information of the library can be embodied in any suitable form, e.g., electronic or biochemical forms. For example, a library of sequence information embodied in electronic form includes an accessible computer data file (or, in biochemical form, a collection of nucleic acid molecules) that contains the representative nucleotide sequences of genes that are differentially expressed (e.g., overexpressed or underexpressed)

as between, for example, i) a cancerous cell and a normal cell; ii) a cancerous cell and a dysplastic cell; iii) a cancerous cell and a cell affected by a disease or condition other than cancer; iv) a metastatic cancerous cell and a normal cell and/or non-metastatic cancerous cell; v) a malignant cancerous cell and a non-malignant cancerous cell (or a normal cell) and/or vi) a dysplastic cell relative to a normal cell. Other combinations and comparisons of cells affected by various diseases or stages of disease will be readily apparent to the ordinarily skilled artisan. Biochemical embodiments of the library include a collection of nucleic acids that have the sequences of the genes in the library, where the nucleic acids can correspond to the entire gene in the library or to a fragment thereof, as described in greater detail below.

The polynucleotide libraries of the subject invention include sequence information of a plurality of polynucleotide sequences, where at least one of the polynucleotides has a sequence of any of "SEQ ID NOS:1-5252." By plurality is meant at least 2, usually at least 3 and can include up to all of "SEQ ID NOS:1-5252." The length and number of polynucleotides in the library will vary with the nature of the library, *e.g.*, if the library is an oligonucleotide array, a cDNA array, a computer database of the sequence information, etc.

Where the library is an electronic library, the nucleic acid sequence information can be present in a variety of media. "Media" refers to a manufacture, other than an isolated nucleic acid molecule, that contains the sequence information of the present invention. Such a manufacture provides the genome sequence or a subset thereof in a form that can be examined by means not directly applicable to the sequence as it exists in a nucleic acid. For example, the nucleotide sequence of the present invention, e.g. the nucleic acid sequences of any of the polynucleotides of "SEQ ID NOS:1-5252," can be recorded on computer readable media, e.g. any medium that can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as a floppy disc, a hard disc storage medium, and a magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. One of skill in the art can readily appreciate how any of the presently known computer readable mediums can be used to create a manufacture comprising a recording of the present sequence information.

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"Recorded" refers to a process for storing information on computer readable medium, using any such methods as known in the art. Any convenient data storage structure can be chosen, based on the means used to access the stored information. A variety of data processor programs and formats can be used for storage, *e.g.* word processing text file, database format, *etc.* In addition to the sequence information, electronic versions of the libraries of the invention can be provided in conjunction or connection with other computer-readable information and/or other types of computer-readable files (*e.g.*, searchable files, executable files, *etc*, including, but not limited to, for example, search program software, *etc.*.).

By providing the nucleotide sequence in computer readable form, the information can be accessed for a variety of purposes. Computer software to access sequence information is publicly available. For example, the BLAST (Altschul *et al.*, *supra.*) and BLAZE (Brutlag *et al.* Comp. Chem. (1993) 17:203) search algorithms on a Sybase system can be used to identify open reading frames (ORFs) within the genome that contain homology to ORFs from other organisms.

As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A skilled artisan can readily appreciate that any one of the currently available computer-based system are suitable for use in the present invention. The data storage means can comprise any manufacture comprising a recording of the present sequence information as described above, or a memory access means that can access such a manufacture.

"Search means" refers to one or more programs implemented on the computer-based system, to compare a target sequence or target structural motif with the stored sequence information. Search means are used to identify fragments or regions of the genome that match a particular target sequence or target motif. A variety of known algorithms are publicly known and commercially available, *e.g.* MacPattern (EMBL), BLASTN and BLASTX (NCBI). A "target sequence" can be any DNA or amino acid

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sequence of six or more nucleotides or two or more amino acids, preferably from about 10 to 100 amino acids or from about 30 to 300 nucleotide residues.

A "target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) are chosen based on a three-dimensional configuration that is formed upon the folding of the target motif, or on consensus sequences of regulatory or active sites. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzyme active sites and signal sequences. Nucleic acid target motifs include, but are not limited to, hairpin structures, promoter sequences and other expression elements such as binding sites for transcription factors.

A variety of structural formats for the input and output means can be used to input and output the information in the computer-based systems of the present invention. One format for an output means ranks fragments of the genome possessing varying degrees of homology to a target sequence or target motif. Such presentation provides a skilled artisan with a ranking of sequences and identifies the degree of sequence similarity contained in the identified fragment.

A variety of comparing means can be used to compare a target sequence or target motif with the data storage means to identify sequence fragments of the genome. A skilled artisan can readily recognize that any one of the publicly available homology search programs can be used as the search means for the computer based systems of the present invention.

As discussed above, the "library" of the invention also encompasses biochemical libraries of the polynucleotides of "SEQ ID NOS:1-5252," e.g., collections of nucleic acids representing the provided polynucleotides. The biochemical libraries can take a variety of forms, e.g., a solution of cDNAs, a pattern of probe nucleic acids stably associated with a surface of a solid support (i.e., an array) and the like. Of particular interest are nucleic acid arrays in which one or more of "SEQ ID NOS:1-5252" is represented on the array. By array is meant a an article of manufacture that has at least a substrate with at least two distinct nucleic acid targets on one of its surfaces, where the number of distinct nucleic acids can be considerably higher, typically being at least 10 nt, usually at least 20 nt and often at least 25 nt. A variety of different array formats have been developed and are known to those of

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skill in the art, including those described in 5,242,974; 5,384,261; 5,405,783; 5,412,087; 5,424,186; 5,429,807; 5,436,327; 5,445,934; 5,472,672; 5,527,681; 5,529,756; 5,545,531; 5,554,501; 5,556,752; 5,561,071; 5,599,895; 5,624,711; 5,639,603; 5,658,734; WO 93/17126; WO 95/11995; WO 95/35505; EP 742287; and EP 799897. The arrays of the subject invention find use in a variety of applications, including gene expression analysis, drug screening, mutation analysis and the like, as disclosed in the above-listed exemplary patent documents.

In addition to the above nucleic acid libraries, analogous libraries of polypeptides are also provided, where the where the polypeptides of the library will represent at least a portion of the polypeptides encoded by "SEQ ID NOS:1-5252."

VII. Utilities

A. <u>Use of Polynucleotide Probes in Mapping, and in Tissue Profiling</u>

Polynucleotide probes, generally comprising at least 12 contiguous nucleotides of a polynucleotide as shown in the Sequence Listing, are used for a variety of purposes, such as chromosome mapping of the polynucleotide and detection of transcription levels. Additional disclosure about preferred regions of the disclosed polynucleotide sequences is found in the Examples. A probe that hybridizes specifically to a polynucleotide disclosed herein should provide a detection signal at least 5-, 10-, or 20-fold higher than the background hybridization provided with other unrelated sequences.

Probes in Detection of Expression Levels. Nucleotide probes are used to detect expression of a gene corresponding to the provided polynucleotide. In Northern blots, mRNA is separated electrophoretically and contacted with a probe. A probe is detected as hybridizing to an mRNA species of a particular size. The amount of hybridization is quantitated to determine relative amounts of expression, for example under a particular condition. Probes are used for in situ hybridization to cells to detect expression. Probes can also be used *in vivo* for diagnostic detection of hybridizing sequences. Probes are typically labeled with a radioactive isotope. Other types of detectable labels can be used such as chromophores, fluors, and enzymes. Other examples of nucleotide hybridization assays are described in WO92/02526 and U.S. Patent No. 5,124,246.

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The Polymerase Chain Reaction (PCR) is another means for detecting small amounts of target nucleic acids (see, e.g., Mullis et al., Meth. Enzymol. (1987) 155:335; U.S. Patent No. 4,683,195; and U.S. Patent No. 4,683,202). Two primer polynucleotides nucleotides hybridize with the target nucleic acids and are used to prime the reaction. The primers can be composed of sequence within or 3' and 5' to the polynucleotides of the Sequence Listing. Alternatively, if the primers are 3' and 5' to these polynucleotides, they need not hybridize to them or the complements. A thermostable polymerase creates copies of target nucleic acids from the primers using the original target nucleic acids as a template. After a large amount of target nucleic acids is generated by the polymerase, it is detected by methods such as Southern blots. When using the Southern blot method, the labeled probe will hybridize to a polynucleotide of the Sequence Listing or complement.

Furthermore, mRNA or cDNA can be detected by traditional blotting techniques described in Sambrook *et al.*, "Molecular Cloning: A Laboratory Manual" (New York, Cold Spring Harbor Laboratory, 1989). mRNA or cDNA generated from mRNA using a polymerase enzyme can be purified and separated using gel electrophoresis. The nucleic acids on the gel are then blotted onto a solid support, such as nitrocellulose. The solid support is exposed to a labeled probe and then washed to remove any unhybridized probe. Next, the duplexes containing the labeled probe are detected. Typically, the probe is labeled with radioactivity.

Mapping. Polynucleotides of the present invention are used to identify a chromosome on which the corresponding gene resides. Such mapping can be useful in identifying the function of the polynucleotide-related gene by its proximity to other genes with known function. Function can also be assigned to the polynucleotide-related gene when particular syndromes or diseases map to the same chromosome. For example, use of polynucleotide probes in identification and quantification of nucleic acid sequence aberrations is described in U.S. Patent No. 5,783,387.

For example, fluorescence in situ hybridization (FISH) on normal metaphase spreads facilitates comparative genomic hybridization to allow total genome assessment of changes in relative copy number of DNA sequences. See Schwartz and Samad, *Curr*.

Opin. Biotechnol (1994) 8:70; Kallionierri et al., Sem. Cancer Biol. (1993) 4:41: Valdes

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et al., Methods in Molecular Biology (1997) 68:1, Boultwood, ed., Human Press, Totowa, NJ.

Polynucleotides are mapped to particular chromosomes using, for example, radiation hybrids or chromosome-specific hybrid panels. See Leach et al., Advances in Genetics, (1995) 33:63-99; Walter et al., Nature Genetics (1994) 7:22; Walter and Goodfellow, Trends in Genetics (1992) 9:352. Panels for radiation hybrid mapping are available from Research Genetics, Inc., Huntsville, Alabama, USA. Databases for markers using various panels are available via the world wide web at http://F/shgc-www.stanford.edu; and http://www-genome.wi.mit.edu/cgi-bin/contig/rhmapper.pl. The statistical program RHMAP can be used to construct a map based on the data from radiation hybridization with a measure of the relative likelihood of one order versus another. RHMAP is available via the world wide web at http://www.sph.umich.edu/group/statgen/software.

In addition, commercial programs are available for identifying regions of chromosomes commonly associated with disease, such as cancer. Polynucleotides based on the polynucleotides of the invention can be used to probe these regions. For example, if through profile searching a provided polynucleotide is identified as corresponding to a gene encoding a kinase, its ability to bind to a cancer-related chromosomal region will suggest its role as a kinase in one or more stages of tumor cell development/growth. Although some experimentation would be required to elucidate the role, the polynucleotide constitutes a new material for isolating a specific protein that has potential for developing a cancer diagnostic or therapeutic.

<u>Tissue Typing or Profiling.</u> Expression of specific mRNA corresponding to the provided polynucleotides can vary in different cell types and can be tissue-specific. This variation of mRNA levels in different cell types can be exploited with nucleic acid probe assays to determine tissue types. For example, PCR, branched DNA probe assays, or blotting techniques utilizing nucleic acid probes substantially identical or complementary to polynucleotides listed in the Sequence Listing can determine the presence or absence of the corresponding cDNA or mRNA.

For example, a metastatic lesion is identified by its developmental organ or tissue source by identifying the expression of a particular marker of that organ or tissue. If a

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polynucleotide is expressed only in a specific tissue type, and a metastatic lesion is found to express that polynucleotide, then the developmental source of the lesion has been identified. Expression of a particular polynucleotide is assayed by detection of either the corresponding mRNA or the protein product. Immunological methods, such as antibody staining, are used to detect a particular protein product. Hybridization methods can be used to detect particular mRNA species, including but not limited to in situ hybridization and Northern blotting.

<u>Use of Polymorphisms.</u> A polynucleotide of the invention will be useful in forensics, genetic analysis, mapping, and diagnostic applications if the corresponding region of a gene is polymorphic in the human population. Particular polymorphic forms of the provided polynucleotides can be used to either identify a sample as deriving from a suspect or rule out the possibility that the sample derives from the suspect. Any means for detecting a polymorphism in a gene are used, including but not limited to electrophoresis of protein polymorphic variants, differential sensitivity to restriction enzyme cleavage, and hybridization to allele-specific probes.

B. Antibody Production

Expression products of a polynucleotide of the invention, the corresponding mRNA or cDNA, or the corresponding complete gene are prepared and used for raising antibodies for experimental, diagnostic, and therapeutic purposes. For polynucleotides to which a corresponding gene has not been assigned, this provides an additional method of identifying the corresponding gene. The polynucleotide or related cDNA is expressed as described above, and antibodies are prepared. These antibodies are specific to an epitope on the polypeptide encoded by the polynucleotide, and can precipitate or bind to the corresponding native protein in a cell or tissue preparation or in a cell-free extract of an in vitro expression system.

Immunogens for raising antibodies are prepared by mixing the polypeptides encoded by the polynucleotides of the present invention with adjuvants. Alternatively, polypeptides are made as fusion proteins to larger immunogenic proteins. Polypeptides are also covalently linked to other larger immunogenic proteins, such as keyhole limpet hemocyanin. Immunogens are typically administered intradernally, subcutaneously, or intramuscularly. Immunogens are administered to experimental animals such as rabbits,

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sheep, and mice, to generate antibodies. Optionally, the animal spleen cells are isolated and fused with myeloma cells to form hybridomas which secrete monoclonal antibodies. Such methods are well known in the art. According to another method known in the art, the selected polynucleotide is administered directly, such as by intramuscular injection, and expressed in vivo. The expressed protein generates a variety of protein-specific immune responses, including production of antibodies, comparable to administration of the protein.

Preparations of polyclonal and monoclonal antibodies specific for polypeptides encoded by a selected polynucleotide are made using standard methods known in the art. The antibodies specifically bind to epitopes present in the polypeptides encoded by polynucleotides disclosed in the Sequence Listing. Typically, at least 6, 8, 10, or 12 contiguous amino acids are required to form an epitope. However, epitopes which involve non-contiguous amino acids may require more, for example at least 15, 25, or 50 amino acids. A short sequence of a polynucleotide may then be unsuitable for use as an epitope to raise antibodies for identifying the corresponding novel protein, because of the potential for cross-reactivity with a known protein. However, the antibodies can be useful for other purposes, particularly if they identify common structural features of a known protein and a novel polypeptide encoded by a polynucleotide of the invention.

Antibodies that specifically bind to human polypeptides encoded by the provided polypeptides should provide a detection signal at least 5-, 10-, or 20-fold higher than a detection signal provided with other proteins when used in Western blots or other immunochemical assays. Preferably, antibodies that specifically polypeptides of the invention do not bind to other proteins in immunochemical assays at detectable levels and can immunoprecipitate the specific polypeptide from solution.

To test for the presence of serum antibodies to the polypeptide of the invention in a human population, human antibodies are purified by methods well known in the art.

Preferably, the antibodies are affinity purified by passing antiserum over a column to which the corresponding selected polypeptide or fusion protein is bound. The bound antibodies can then be eluted from the column, for example using a buffer with a high salt concentration.

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In addition to the antibodies discussed above, genetically engineered antibody derivatives are made, such as single chain antibodies, according to methods well known in the art.

C. <u>Use of Polynucleotides to Construct Arrays for Diagnostics</u>

Polynucleotide arrays provide a high throughput technique that can assay a large number of polynucleotide sequences in a sample. This technology can be used as a diagnostic and as a tool to test for differential expression to determine function of an encoded protein. Arrays can be created by spotting polynucleotide probes onto a substrate (e.g., glass, nitrocelllose, etc.) in a two-dimensional matrix or array having bound probes.

The probes can be bound to the substrate by either covalent bonds or by non-specific interactions, such as hydrophobic interactions. Samples of polynucleotides can be detectably labeled (*e.g.*, using radioactive or fluorescent labels) and then hybridized to the probes. Double stranded polynucleotides, comprising the labeled sample polynucleotides bound to probe polynucleotides, can be detected once the unbound portion of the sample is washed away. Techniques for constructing arrays and methods of using these arrays are described in EP No. 0 799 897; PCT No. WO 97/29212; PCT No. WO 97/27317; EP No. 0 785 280; PCT No. WO 97/02357; U.S. Pat. No. 5,593,839; U.S. Pat. No. 5,578,832; EP No. 0 728 520; U.S. Pat. No. 5,599,695; EP No. 0 721 016; U.S. Pat. No. 5,556,752; PCT No. WO 95/22058; and U.S. Pat. No. 5,631,734.

As discussed in some detail above, arrays can be used to examine differential expression of genes and can be used to determine gene function. For example, arrays of the instant polynucleotide sequences can be used to determine if any of the provided polynucleotides are differentially expressed between a test cell and control cell (e.g., cancer cells and normal cells). For example, high expression of a particular message in a cancer cell, which is not observed in a corresponding normal cell, can indicate a cancer specific protein. Exemplary uses of arrays are further described in, for example, Pappalarado et al., Sem. Radiation Oncol. (1998) 8:217; and Ramsay Nature Biotechnol. (1998) 16:40.

D. Differential Expression

The polynucleotides of the invention can also be used to detect differences in expression levels between two cells, e.g., as a method to identify abnormal or diseased tissue in a human. For polynucleotides corresponding to profiles of protein families, the

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choice of tissue can be selected according to the putative biological function. In general, the expression of a gene corresponding to a specific polynucleotide is compared between a first tissue that is suspected of being diseased and a second, normal tissue of the human. The tissue suspected of being abnormal or diseased can be derived from a different tissue type of the human, but preferably it is derived from the same tissue type; for example an intestinal polyp or other abnormal growth should be compared with normal intestinal tissue. The normal tissue can be the same tissue as that of the test sample, or any normal tissue of the patient, especially those that express the polynucleotide-related gene of interest (e.g., brain, thymus, testis, heart, prostate, placenta, spleen, small intestine, skeletal muscle, pancreas, and the mucosal lining of the colon). A difference between the polynucleotide-related gene, mRNA, or protein in the two tissues which are compared, for example in molecular weight, amino acid or nucleotide sequence, or relative abundance, indicates a change in the gene, or a gene which regulates it, in the tissue of the human that was suspected of being diseased. Examples of detection of differential expression and its use in diagnosis of cancer are described in U.S. Patent Nos. 5,688,641 and 5,677,125.

The polynucleotide-related genes in the two tissues are compared by any means known in the art. For example, the two genes can be sequenced, and the sequence of the gene in the tissue suspected of being diseased compared with the gene sequence in the normal tissue. The genes corresponding to a provided polynucleotide, or portions thereof, in the two tissues are amplified, for example using nucleotide primers based on the nucleotide sequence shown in the Sequence Listing, using the polymerase chain reaction. The amplified genes or portions of genes are hybridized to detectably labeled nucleotide probes selected from a nucleotide sequence shown in the Sequence Listing. A difference in the nucleotide sequence of the isolated gene in the tissue suspected of being diseased compared with the normal nucleotide sequence suggests a role of the gene product encoded by the subject polynucleotide in the disease, and provides guidance for preparing a therapeutic agent.

Alternatively, mRNA corresponding to a provided polynucleotide in the two tissues is compared. PolyA⁺RNA is isolated from the two tissues as is known in the art. For example, one of skill in the art can readily determine differences in the size or amount of mRNA transcripts between the two tissues using Northern blots and detectably labeled

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nucleotide probes selected from the nucleotide sequence shown in the Sequence Listing. Increased or decreased expression of a given mRNA in a tissue sample suspected of being diseased, compared with the expression of the same mRNA in a normal tissue, suggests that the expressed protein has a role in the disease, and also provides a lead for preparing a therapeutic agent.

The comparison can also be accomplished by analyzing polypeptides between the matched samples. The sizes of the proteins in the two tissues are compared, for example, using antibodies of the present invention to detect polypeptides in Western blots of protein extracts from the two tissues. Other changes, such as expression levels and subcellular localization, can also be detected immunologically, using antibodies to the corresponding protein. A higher or lower level of expression of a given polypeptide in a tissue suspected of being diseased, compared with the same protein expression level in a normal tissue, is indicative that the expressed protein has a role in the disease, and provides guidance for preparing a therapeutic agent.

Similarly, comparison of polynucleotide sequences or of gene expression products, e.g., mRNA and protein, between a human tissue that is suspected of being diseased and a normal tissue of a human, are used to follow disease progression or remission in the human. Such comparisons are made as described above. For example, increased or decreased expression of a gene corresponding to an inventive polynucleotide in the tissue suspected of being neoplastic can indicate the presence of neoplastic cells in the tissue. The degree of increased expression of a given gene in the neoplastic tissue relative to expression of the same gene in normal tissue, or differences in the amount of increased expression of a given gene in the neoplastic tissue over time, is used to assess the progression of the neoplasia in that tissue or to monitor the response of the neoplastic tissue to a therapeutic protocol over time.

The expression pattern of any two cell types can be compared, such as low and high metastatic tumor cell lines, malignant or non-malignant cells, or cells from tissue which have and have not been exposed to a therapeutic agent. A genetic predisposition to disease in a human is detected by comparing expression levels of an mRNA or protein corresponding to a polynucleotide of the invention in a fetal tissue with levels associated in normal fetal tissue. Fetal tissues that are used for this purpose include, but are not limited

to, amniotic fluid, chorionic villi, blood, and the blastomere of an in vitro-fertilized embryo. The comparable normal polynucleotide-related gene is obtained from any tissue. The mRNA or protein is obtained from a normal tissue of a human in which the polynucleotide-related gene is expressed. Differences such as alterations in the nucleotide sequence or size of the same product of the fetal polynucleotide-related gene or mRNA, or alterations in the molecular weight, amino acid sequence, or relative abundance of fetal protein, can indicate a germline mutation in the polynucleotide-related gene of the fetus, which indicates a genetic predisposition to disease. Particular diagnostic and prognostic uses of the disclosed polynucleotides are described in more detail below.

E. Diagnostic, Prognostic, and Other Uses Based On Differential Expression

In general, diagnostic methods of the invention for involve detection of a level or amount of a gene product, particularly a differentially expressed gene product, in a test sample obtained from a patient suspected of having or being susceptible to a disease (e.g., breast cancer, lung cancer, colon cancer and/or metastatic forms thereof), and comparing the detected levels to those levels found in normal cells (e.g., cells substantially unaffected by cancer) and/or other control cells (e.g., to differentiate a cancerous cell from a cell affected by dysplasia). Furthermore, the severity of the disease can be assessed by comparing the detected levels of a differentially expressed gene product with those levels detected in samples representing the levels of differentially gene product associated with varying degrees of severity of disease.

The term "differentially expressed gene" is intended to encompass a polynucleotide that can, for example, include an open reading frame encoding a gene product (e.g., a polypeptide), and/or introns of such genes and adjacent 5' and 3' non-coding nucleotide sequences involved in the regulation of expression, up to about 20 kb beyond the coding region, but possibly further in either direction. The gene can be introduced into an appropriate vector for extrachromosomal maintenance or for integration into a host genome. In general, a difference in expression level associated with a decrease in expression level of at least about 25%, usually at least about 50% to 75%, more usually at least about 90% or more is indicative of a differentially expressed gene of interest, i.e., a gene that is underexpressed or down-regulated in the test sample relative to a control sample. Furthermore, a difference in expression level associated with an increase in

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expression of at least about 25%, usually at least about 50% to 75%, more usually at least about 90% and can be at least about 1½-fold, usually at least about 2-fold to about 10-fold, and can be about 100-fold to about 1,000-fold increase relative to a control sample is indicative of a differentially expressed gene of interest, *i.e.*, an overexpressed or upregulated gene.

"Differentially expressed polynucleotide" as used herein means a nucleic acid molecule (RNA or DNA) having a sequence that represents a differentially expressed gene, e.g., the differentially expressed polynucleotide comprises a sequence (e.g., an open reading frame encoding a gene product) that uniquely identifies a differentially expressed gene so that detection of the differentially expressed polynucleotide in a sample is correlated with the presence of a differentially expressed gene in a sample. "Differentially expressed polynucleotides" is also meant to encompass fragments of the disclosed polynucleotides, e.g., fragments retaining biological activity, as well as nucleic acids homologous, substantially similar, or substantially identical (e.g., having about 90% sequence identity) to the disclosed polynucleotides.

Methods of the subject invention useful in diagnosis or prognosis typically involve comparison of the abundance of a selected differentially expressed gene product in a sample of interest with that of a control to determine any relative differences in the expression of the gene product, where the difference can be measured qualitatively and/or quantitatively. Quantitation can be accomplished, for example, by comparing the level of expression product detected in the sample with the amounts of product present in a standard curve. A comparison can be made visually; by using a technique such as densitometry, with or without computerized assistance; by preparing a representative library of cDNA clones of mRNA isolated from a test sample, sequencing the clones in the library to determine that number of cDNA clones corresponding to the same gene product, and analyzing the number of clones corresponding to that same gene product relative to the number of clones of the same gene product in a control sample; or by using an array to detect relative levels of hybridization to a selected sequence or set of sequences, and comparing the hybridization pattern to that of a control. The differences in expression are then correlated with the presence or absence of an abnormal expression pattern. A variety of different methods for determining the nucleic acid abundance in a sample are known to

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those of skill in the art, where particular methods of interest include those described in:
Pietu et al. Genome Res. (1996) 6:492; Zhao et al., Gene (1995) 156:207; Soares, Curr.

Opin. Biotechnol. (1977) 8: 542; Raval, J. Pharmacol Toxicol Methods (1994) 32:125;

Chalifour et al., Anal. Biochem (1994) 216:299; Stolz et al., Mol. Biotechnol. (1996) 6:225;

Hong et al., Biosci. Reports (1982) 2:907; and McGraw, Anal. Biochem. (1984) 143:298.

Also of interest are the methods disclosed in WO 97/27317, the disclosure of which is herein incorporated by reference.

In general, diagnostic assays of the invention involve detection of a gene product of a the polynucleotide sequence (e.g., mRNA or polypeptide) that corresponds to a sequence of "SEQ ID NOS:1-5252." The patient from whom the sample is obtained can be apparently healthy, susceptible to disease (e.g., as determined by family history or exposure to certain environmental factors), or can already be identified as having a condition in which altered expression of a gene product of the invention is implicated.

In the assays of the invention, the diagnosis can be determined based on detected gene product expression levels of a gene product encoded by at least one, preferably at least two or more, at least 3 or more, or at least 4 or more of the polynucleotides having a sequence set forth in "SEQ ID NOS:1-5252," and can involve detection of expression of genes corresponding to all of "SEQ ID NOS:1-5252" and/or additional sequences that can serve as additional diagnostic markers and/or reference sequences. Where the diagnostic method is designed to detect the presence or susceptibility of a patient to cancer, the assay preferably involves detection of a gene product encoded by a gene corresponding to a polynucleotide that is differentially expressed in cancer. For example, a higher level of expression of a polynucleotide corresponding to SEQ ID NO:2024 relative to a level associated with a normal sample can indicate the presence of cancer in the patient from whom the sample is derived. In another example, detection of a lower level of a polynucleotide corresponding to SEQ ID NO:590 relative to a normal level is indicative of the presence of cancer in the patient. Further examples of such differentially expressed polynucleotides are described in the Examples below. Given the provided polynucleotides and information regarding their relative expression levels provided herein, assays using such polynucleotides and detection of their expression levels in diagnosis and prognosis will be readily apparent to the ordinarily skilled artisan.

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Any of a variety of detectable labels can be used in connection with the various embodiments of the diagnostic methods of the invention. Suitable detectable labels include fluorochromes, (e.g. fluorescein isothiocyanate (FITC), rhodamine, Texas Red, phycoerythrin, allophycocyanin, 6-carboxyfluorescein (6-FAM), 2',7'-dimethoxy-4',5'-dichloro-6-carboxyfluorescein, 6-carboxy-X-rhodamine (ROX), 6-carboxy-2',4',7',4,7-hexachlorofluorescein (HEX), 5-carboxyfluorescein (5-FAM) or N,N,N',N'-tetramethyl-6-carboxyrhodamine (TAMRA)), radioactive labels, (e.g. ³²P, ³⁵S, ³H, etc.), and the like. The detectable label can involve a two stage systems (e.g., biotin-avidin, hapten-anti-hapten antibody, etc.)

Reagents specific for the polynucleotides and polypeptides of the invention, such as antibodies and nucleotide probes, can be supplied in a kit for detecting the presence of an expression product in a biological sample. The kit can also contain buffers or labeling components, as well as instructions for using the reagents to detect and quantify expression products in the biological sample. Exemplary embodiments of the diagnostic methods of the invention are described below in more detail.

Polypeptide detection in diagnosis. In one embodiment, the test sample is assayed for the level of a differentially expressed polypeptide. Diagnosis can be accomplished using any of a number of methods to determine the absence or presence or altered amounts of the differentially expressed polypeptide in the test sample. For example, detection can utilize staining of cells or histological sections with labeled antibodies, performed in accordance with conventional methods. Cells can be permeabilized to stain cytoplasmic molecules. In general, antibodies that specifically bind a differentially expressed polypeptide of the invention are added to a sample, and incubated for a period of time sufficient to allow binding to the epitope, usually at least about 10 minutes. The antibody can be detectably labeled for direct detection (e.g., using radioisotopes, enzymes, fluorescers, chemiluminescers, and the like), or can be used in conjunction with a second stage antibody or reagent to detect binding (e.g., biotin with horseradish peroxidaseconjugated avidin, a secondary antibody conjugated to a fluorescent compound, e.g. fluorescein, rhodamine, Texas red, etc.). The absence or presence of antibody binding can be determined by various methods, including flow cytometry of dissociated cells, microscopy, radiography, scintillation counting, etc. Any suitable alternative methods can

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of qualitative or quantitative detection of levels or amounts of differentially expressed polypeptide can be used, for example ELISA, western blot, immunoprecipitation, radioimmunoassay, etc.

In general, the detected level of differentially expressed polypeptide in the test sample is compared to a level of the differentially expressed gene product in a reference or control sample, *e.g.*, in a normal cell (negative control) or in a cell having a known disease state (positive control).

mRNA detection. The diagnostic methods of the invention can also or alternatively involve detection of mRNA encoded by a gene corresponding to a differentially expressed polynucleotides of the invention. Any suitable qualitative or quantitative methods known in the art for detecting specific mRNAs can be used. mRNA can be detected by, for example, *in situ* hybridization in tissue sections, by reverse transcriptase-PCR, or in Northern blots containing poly A+ mRNA. One of skill in the art can readily use these methods to determine differences in the size or amount of mRNA transcripts between two samples. For example, the level of mRNA of the invention in a tissue sample suspected of being cancerous or dysplastic is compared with the expression of the mRNA in a reference sample, *e.g.*, a positive or negative control sample (*e.g.*, normal tissue, cancerous tissue, *etc.*).

Any suitable method for detecting and comparing mRNA expression levels in a sample can be used in connection with the diagnostic methods of the invention (see, e.g., U.S. 5,804,382). For example, mRNA expression levels in a sample can be determined by generation of a library of expressed sequence tags (ESTs) from the sample, where the EST library is representative of sequences present in the sample (Adams, et al., (1991) Science 252:1651). Enumeration of the relative representation of ESTs within the library can be used to approximate the relative representation of the gene transcript within the starting sample. The results of EST analysis of a test sample can then be compared to EST analysis of a reference sample to determine the relative expression levels of a selected polynucleotide, particularly a polynucleotide corresponding to one or more of the differentially expressed genes described herein.

Alternatively, gene expression in a test sample can be performed using serial analysis of gene expression (SAGE) methodology (Velculescu et al., *Science* (1995)

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270:484). In short, SAGE involves the isolation of short unique sequence tags from a specific location within each transcript. The sequence tags are concatenated, cloned, and sequenced. The frequency of particular transcripts within the starting sample is reflected by the number of times the associated sequence tag is encountered with the sequence population.

Gene expression in a test sample can also be analyzed using differential display (DD) methodology. In DD, fragments defined by specific sequence delimiters (e.g., restriction enzyme sites) are used as unique identifiers of genes, coupled with information about fragment length or fragment location within the expressed gene. The relative representation of an expressed gene with a sample can then be estimated based on the relative representation of the fragment associated with that gene within the pool of all possible fragments. Methods and compositions for carrying out DD are well known in the art, see, e.g., U.S. 5,776,683; and U.S. 5,807,680.

Alternatively, gene expression in a sample using hybridization analysis, which is based on the specificity of nucleotide interactions. Oligonucleotides or cDNA can be used to selectively identify or capture DNA or RNA of specific sequence composition, and the amount of RNA or cDNA hybridized to a known capture sequence determined qualitatively or quantitatively, to provide information about the relative representation of a particular message within the pool of cellular messages in a sample. Hybridization analysis can be designed to allow for concurrent screening of the relative expression of hundreds to thousands of genes by using, for example, array-based technologies having high density formats, including filters, microscope slides, or microchips, or solution-based technologies that use spectroscopic analysis (e.g., mass spectrometry). One exemplary use of arrays in the diagnostic methods of the invention is described below in more detail.

Use of a single gene in diagnostic applications. The diagnostic methods of the invention can focus on the expression of a single differentially expressed gene. For example, the diagnostic method can involve detecting a differentially expressed gene, or a polymorphism of such a gene (e.g., a polymorphism in an coding region or control region), that is associated with disease. Disease-associated polymorphisms can include deletion or truncation of the gene, mutations that alter expression level and/or affect activity of the encoded protein, etc.

Changes in the promoter or enhancer sequence that affect expression levels of an differentially gene can be compared to expression levels of the normal allele by various methods known in the art. Methods for determining promoter or enhancer strength include quantitation of the expressed natural protein; insertion of the variant control element into a vector with a reporter gene such as β -galactosidase, luciferase, chloramphenicol acetyltransferase, *etc.* that provides for convenient quantitation; and the like.

A number of methods are available for analyzing nucleic acids for the presence of a specific sequence, e.g. a disease associated polymorphism. Where large amounts of DNA are available, genomic DNA is used directly. Alternatively, the region of interest is cloned into a suitable vector and grown in sufficient quantity for analysis. Cells that express a differentially expressed gene can be used as a source of mRNA, which can be assayed directly or reverse transcribed into cDNA for analysis. The nucleic acid can be amplified by conventional techniques, such as the polymerase chain reaction (PCR), to provide sufficient amounts for analysis, and a detectable label can be included in the amplification reaction (e.g., using a detectably labeled primer or detectably labeled oligonucleotides) to facilitate detection. The use of the polymerase chain reaction is described in Saiki, et al., Science (1985) 239:487, and a review of techniques can be found in Sambrook, et al., Molecular Cloning: A Laboratory Manual, (1989) pp. 14.2. Alternatively, various methods are known in the art that utilize oligonucleotide ligation as a means of detecting polymorphisms, for examples see Riley et al., Nucl. Acids Res. (1990) 18:2887; and Delahunty et al., Am. J. Hum. Genet. (1996) 58:1239.

The sample nucleic acid, e.g. amplified or cloned fragment, is analyzed by one of a number of methods known in the art. The nucleic acid can be sequenced by dideoxy or other methods, and the sequence of bases compared to a selected sequence, e.g., to a wild-

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type sequence. Hybridization with the polymorphic or variant sequence can also be used to determine its presence in a sample (e.g., by Southern blot, dot blot, etc.). The hybridization pattern of a polymorphic or variant sequence and a control sequence to an array of oligonucleotide probes immobilized on a solid support, as described in US 5,445,934, or in WO 95/35505, can also be used as a means of identifying polymorphic or variant sequences associated with disease. Single strand conformational polymorphism (SSCP) analysis, denaturing gradient gel electrophoresis (DGGE), and heteroduplex analysis in gel matrices are used to detect conformational changes created by DNA sequence variation as alterations in electrophoretic mobility. Alternatively, where a polymorphism creates or destroys a recognition site for a restriction endonuclease, the sample is digested with that endonuclease, and the products size fractionated to determine whether the fragment was digested. Fractionation is performed by gel or capillary electrophoresis, particularly acrylamide or agarose gels.

Screening for mutations in an differentially expressed gene can be based on the functional or antigenic characteristics of the protein. Protein truncation assays are useful in detecting deletions that can affect the biological activity of the protein. Various immunoassays designed to detect polymorphisms in proteins can be used in screening. Where many diverse genetic mutations lead to a particular disease phenotype, functional protein assays have proven to be effective screening tools. The activity of the encoded protein can be determined by comparison with the wild-type protein.

Pattern matching in diagnosis using arrays. In another embodiment, the diagnostic and/or prognostic methods of the invention involve detection of expression of a selected set of genes in a test sample to produce a test expression pattern (TEP). The TEP is compared to a reference expression pattern (REP), which is generated by detection of expression of the selected set of genes in a reference sample (e.g., a positive or negative control sample). The selected set of genes includes at least one of the genes of the invention, which genes correspond to the polynucleotide sequences of "SEQ ID NOS:1-5252." Of particular interest is a selected set of genes that includes gene differentially expressed in the disease for which the test sample is to be screened.

"Reference sequences" or "reference polynucleotides" as used herein in the context of differential gene expression analysis and diagnosis/prognosis refers to a selected set of

polynucleotides, which selected set includes at least one or more of the differentially expressed polynucleotides described herein. A plurality of reference sequences, preferably comprising positive and negative control sequences, can be included as reference sequences. Additional suitable reference sequences are found in Genbank, Unigene, and other nucleotide sequence databases (including, *e.g.*, expressed sequence tag (EST), partial, and full-length sequences).

"Reference array" means an array having reference sequences for use in hybridization with a sample, where the reference sequences include all, at least one of, or any subset of the differentially expressed polynucleotides described herein. Usually such an array will include at least 3 different reference sequences, and can include any one or all of the provided differentially expressed sequences. Arrays of interest can further comprise sequences, including polymorphisms, of other genetic sequences, particularly other sequences of interest for screening for a disease or disorder (e.g., cancer, dysplasia, or other related or unrelated diseases, disorders, or conditions). The oligonucleotide sequence on the array will usually be at least about 12 nt in length, and can be of about the length of the provided sequences, or can extend into the flanking regions to generate fragments of 100 nt to 200 nt in length or more.

A "reference expression pattern" or "REP" as used herein refers to the relative levels of expression of a selected set of genes, particularly of differentially expressed genes, that is associated with a selected cell type, e.g., a normal cell, a cancerous cell, a cell exposed to an environmental stimulus, and the like. A "test expression pattern" or "TEP" refers to relative levels of expression of a selected set of genes, particularly of differentially expressed genes, in a test sample (e.g., a cell of unknown or suspected disease state, from which mRNA is isolated).

"Diagnosis" as used herein generally includes determination of a subject's susceptibility to a disease or disorder, determination as to whether a subject is presently affected by a disease or disorder, as well as to the prognosis of a subject affected by a disease or disorder (e.g., identification of pre-metastatic or metastatic cancerous states, stages of cancer, or responsiveness of cancer to therapy). The present invention particularly encompasses diagnosis of subjects in the context of breast cancer (e.g., carcinoma in situ (e.g., ductal carcinoma in situ), estrogen receptor (ER)-positive breast

cancer, ER-negative breast cancer, or other forms and/or stages of breast cancer), lung cancer (e.g., small cell carcinoma, non-small cell carcinoma, mesothelioma, and other forms and/or stages of lung cancer), and colon cancer (e.g., adenomatous polyp, colorectal carcinoma, and other forms and/or stages of colon cancer).

"Sample" or "biological sample" as used throughout here are generally meant to refer to samples of biological fluids or tissues, particularly samples obtained from tissues, especially from cells of the type associated with the disease for which the diagnostic application is designed (e.g., ductal adenocarcinoma), and the like. "Samples" is also meant to encompass derivatives and fractions of such samples (e.g., cell lysates). Where the sample is solid tissue, the cells of the tissue can be dissociated or tissue sections can be analyzed.

REPs can be generated in a variety of ways according to methods well known in the art. For example, REPs can be generated by hybridizing a control sample to an array having a selected set of polynucleotides (particularly a selected set of differentially expressed polynucleotides), acquiring the hybridization data from the array, and storing the data in a format that allows for ready comparison of the REP with a TEP. Alternatively, all expressed sequences in a control sample can be isolated and sequenced, *e.g.*, by isolating mRNA from a control sample, converting the mRNA into cDNA, and sequencing the cDNA. The resulting sequence information roughly or precisely reflects the identity and relative number of expressed sequences in the sample. The sequence information can then be stored in a format (*e.g.*, a computer-readable format) that allows for ready comparison of the REP with a TEP. The REP can be normalized prior to or after data storage, and/or can be processed to selectively remove sequences of expressed genes that are of less interest or that might complicate analysis (*e.g.*, some or all of the sequences associated with housekeeping genes can be eliminated from REP data).

TEPs can be generated in a manner similar to REPs, e.g., by hybridizing a test sample to an array having a selected set of polynucleotides, particularly a selected set of differentially expressed polynucleotides, acquiring the hybridization data from the array, and storing the data in a format that allows for ready comparison of the TEP with a REP. The REP and TEP to be used in a comparison can be generated simultaneously, or the TEP can be compared to previously generated and stored REPs.

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In one embodiment of the invention, comparison of a TEP with a REP involves hybridizing a test sample with a reference array, where the reference array has one or more reference sequences for use in hybridization with a sample. The reference sequences include all, at least one of, or any subset of the differentially expressed polynucleotides described herein. Hybridization data for the test sample is acquired, the data normalized, and the produced TEP compared with a REP generated using an array having the same or similar selected set of differentially expressed polynucleotides. Probes that correspond to sequences differentially expressed between the two samples will show decreased or increased hybridization efficiency for one of the samples relative to the other.

Reference arrays can be produced according to any suitable methods known in the art. For example, methods of producing large arrays of oligonucleotides are described in U.S. 5,134,854, and U.S. 5,445,934 using light-directed synthesis techniques. Using a computer controlled system, a heterogeneous array of monomers is converted, through simultaneous coupling at a number of reaction sites, into a heterogeneous array of polymers. Alternatively, microarrays are generated by deposition of pre-synthesized oligonucleotides onto a solid substrate, for example as described in PCT published application no. WO 95/35505.

Methods for collection of data from hybridization of samples with a reference arrays are also well known in the art. For example, the polynucleotides of the reference and test samples can be generated using a detectable fluorescent label, and hybridization of the polynucleotides in the samples detected by scanning the microarrays for the presence of the detectable label. Methods and devices for detecting fluorescently marked targets on devices are known in the art. Generally, such detection devices include a microscope and light source for directing light at a substrate. A photon counter detects fluorescence from the substrate, while an x-y translation stage varies the location of the substrate. A confocal detection device that can be used in the subject methods is described in U.S. Patent no. 5,631,734. A scanning laser microscope is described in Shalon et al., *Genome Res.* (1996) 6:639. A scan, using the appropriate excitation line, is performed for each fluorophore used. The digital images generated from the scan are then combined for subsequent analysis. For any particular array element, the ratio of the fluorescent signal from one

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sample (e.g., a test sample) is compared to the fluorescent signal from another sample (e.g., a reference sample), and the relative signal intensity determined.

Methods for analyzing the data collected from hybridization to arrays are well known in the art. For example, where detection of hybridization involves a fluorescent label, data analysis can include the steps of determining fluorescent intensity as a function of substrate position from the data collected, removing outliers, *i.e.* data deviating from a predetermined statistical distribution, and calculating the relative binding affinity of the targets from the remaining data. The resulting data can be displayed as an image with the intensity in each region varying according to the binding affinity between targets and probes.

In general, the test sample is classified as having a gene expression profile corresponding to that associated with a disease or non-disease state by comparing the TEP generated from the test sample to one or more REPs generated from reference samples (e.g., from samples associated with cancer or specific stages of cancer, dysplasia, samples affected by a disease other than cancer, normal samples, etc.). The criteria for a match or a substantial match between a TEP and a REP include expression of the same or substantially the same set of reference genes, as well as expression of these reference genes at substantially the same levels (e.g., no significant difference between the samples for a signal associated with a selected reference sequence after normalization of the samples, or at least no greater than about 25% to about 40% difference in signal strength for a given reference sequence. In general, a pattern match between a TEP and a REP includes a match in expression, preferably a match in qualitative or quantitative expression level, of at least one of, all or any subset of the differentially expressed genes of the invention.

Pattern matching can be performed manually, or can be performed using a computer program. Methods for preparation of substrate matrices (*e.g.*, arrays), design of oligonucleotides for use with such matrices, labeling of probes, hybridization conditions, scanning of hybridized matrices, and analysis of patterns generated, including comparison analysis, are described in, for example, U.S. 5,800,992.

F. Use of the Polynucleotides of the Invention in Cancer

Oncogenesis involves the unbridled growth, dedifferentiation and abnormal migration of cells. Cancerous cells can have the ability to compress, invade, and destroy

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normal tissue. Cancerous cells may also metastasize to other parts of the body via the bloodstream or the lymph system and colonize in these other areas. Different cancers are classified by the cell from which the cancerous cell is derived and from its cellular morphology and/or state of differentiation.

Somatic genetic abnormalities cause cancer initiation and progression. Cancer generally is clonally formed, *i.e.* gain of function of oncogenes and loss of function of tumor suppressor genes within a single cell transform the cell to be cancerous, and that single cell grows and divides to form a cancerous lesion. The genes known to be involved in cancer initiation and progression are involved in numerous cellular functions, including developmental differentiation, cell cycle regulation, cell signaling, immunological response, DNA replication, and DNA repair.

The identification and characterization of genetic or biochemical markers in blood or tissues that will detect the earliest changes along the carcinogenesis pathway and monitor the efficacy of various therapies and preventive interventions is a major goal of cancer research. Scientists have identified genetic changes in stool specimens that indicate the stages of colon cancer, and other biomarkers such as gene mutations, hormone receptors, proteins that inhibit metastasis, and enzymes that metabolize drugs are all being used to determine the severity and predict the course of breast, prostate, lung, and other cancers.

Recent advances in the pathogenesis of certain cancers has been helpful in determining patient treatment. The level of expression of certain polynucleotides can be indicative of a poorer prognosis, and therefore warrant more aggressive chemo- or radio-therapy for a patient. The correlation of novel surrogate tumor specific features with response to treatment and outcome in patients has defined certain prognostic indicators that allow the design of tailored therapy based on the molecular profile of the tumor. These therapies include antibody targeting and gene therapy. Moreover, a promising level of one or more marker polynucleotides can provide impetus for not aggressively treating a particular patient, thus sparing the patient the deleterious side effects of aggressive therapy. Determining expression of certain polynucleotides and comparison of a patients profile with known expression in normal tissue and variants of the disease allows

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a determination of the best possible treatment for a patient, both in terms of specificity of treatment and in terms of comfort level of the patient.

Surrogate tumor markers, such as polynucleotide expression, can also be used to better classify, and thus diagnose and treat, different forms and disease states of cancer. Two classifications widely used in oncology that can benefit from identification of the expression levels of the polynucleotides of the invention are staging of the cancerous disorder, and grading the nature of the cancerous tissue.

Staging. Staging is a process used by physicians to describe how advanced the cancerous state is in a patient. Staging assists the physician in determining a prognosis, planning treatment and evaluating the results of such treatment. Different staging systems are used for different types of cancer, but each generally involves the following determinations: the type of tumor, indicated by T; whether the cancer has metastasized to nearby lymph nodes, indicated by N; and whether the cancer has metastasized to more distant parts of the body, indicated by M. This system of staging is called the TNM system. Generally, if a cancer is only detectable in the area of the primary lesion without having spread to any lymph nodes it is called Stage I. If it has spread only to the closest lymph nodes, it is called Stage II. In Stage III, the cancer has generally spread to the lymph nodes in near proximity to the site of the primary lesion. Cancers that have spread to a distant part of the body, such as the liver, bone, brain or another site, are called Stage IV, the most advanced stage.

Currently, the determination of staging is done using pathological techniques and is based more on the presence or absence of malignant tissue rather than the characteristics of the tumor type. Presence or absence of malignant tissue is based primarily on the gross morphology of the cells in the areas biopsied. The polynucleotides of the invention can facilitate fine-tuning of the staging process by identifying markers for the aggresivity of a cancer, e.g. the metastatic potential, as well as the presence in different areas of the body. Thus, a Stage II cancer with a polynucleotide signifying a high metastatic potential cancer can be used to change a borderline Stage II tumor to a Stage III tumor, justifying more aggressive therapy. Conversely, the presence of a polynucleotide signifying a lower metastatic potential allows more conservative staging of a tumor.

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Grading of cancers. Grade is a term used to describe how closely a tumor resembles normal tissue of its same type. Based on the microscopic appearance of a tumor, pathologists will identify the grade of a tumor based on parameters such as cell morphology, cellular organization, and other markers of differentiation. As a general rule, the grade of a tumor corresponds to its rate of growth or aggressiveness. That is, undifferentiated or high-grade tumors grow more quickly than well differentiated or low-grade tumors. Information about tumor grade is useful in planning treatment and predicting prognosis.

The American Joint Commission on Cancer has recommended the following guidelines for grading tumors: 1) GX Grade cannot be assessed; 2) G1 Well differentiated; G2 Moderately well differentiated; 3) G3 Poorly differentiated; 4) G4 Undifferentiated. Although grading is used by pathologists to describe most cancers, it plays a more important role in treatment planning for certain types than for others. An example is the Gleason system that is specific for prostate cancer, which uses grade numbers to describe the degree of differentiation. Lower Gleason scores indicate well-differentiated cells. Intermediate scores denote tumors with moderately differentiated cells. Higher scores describe poorly differentiated cells. Grade is also important in some types of brain tumors and soft tissue sarcomas.

The polynucleotides of the invention can be especially valuable in determining the grade of the tumor, as they not only can aid in determining the differentiation status of the cells of a tumor, they can also identify factors other than differentiation that are valuable in determining the aggressivity of a tumor, such as metastatic potential.

<u>Familial Cancer Genes.</u> A number of cancer syndromes are linked to Mendelian inheritance of a predisposition to develop particular cancers. The following table contains a list of cancer types that can be inherited, and for which the gene or genes responsible have been identified. Most of the cancer types listed can occur as part of several different genetic conditions, each caused by alterations in a different gene.

Cancer Type	Genetic Condition	Gene
Brain	Li-Fraumeni syndrome	TP53
Brain	Neurofibromatosis!	NF1
	Neurofibromatosis 2	NF2
	von Hippel-Lindau syndrome	VHL

Cancer Type	Genetic Condition	Gene
-	Tuberous sclerosis 2	TSC2
Breast	Hereditary breast/ovarian cancer 1	BRCA1
	Hereditary breast/ovarian cancer 2	BRCA2
	Li-Fraumeni syndrome	TP53
	Ataxia telangiectasia	ATM
Colon	Familial adenomatous polyposis (FAP)	APC
	Hereditary non-polyposis colon cancer (HNPCC) 1	HMSH2
	Hereditary non-polyposis colon cancer (HNPCC) 2	hMLH1
	Hereditary non-polyposis colon cancer (HNPCC) 3	hPMS1
	Hereditary non-polyposis colon cancer (HNPCC) 4	hPMS2
Endocrine	Multiple endocrine neoplasia 1 (MEN1)	MEN1
(parathyroid, pituitary, GI endocrine)	• • • • •	
Endocrine	Multiple endocrine neoplasia 2 (MEN2)	RET
(pheochromacytoma, medullary thyroid)		
Endometrial	Hereditary non-polyposis colon cancer (HNPCC) 1	hMSH2
	Hereditary non-polyposis colon cancer (HNPCC) 2	hMLH1
	Hereditary non-polyposis colon cancer (HNPCC) 3	hPMS1
	Hereditary non-polyposis colon cancer (HNPCC) 4	hPMS2
Eye	Hereditary retinoblastoma	RB1
Hematologic	Li-Fraumeni syndrome	TP53
(lymphomas and leukemia)		
	Ataxia telangiectasia	ATM
Kidney	Hereditary Wilms' tumor	WT1
•	von Hippel-Lindau syndrome	VHL
	Tuberous sclerosis 2	TSC2
Ovary	Hereditary breast/ovarian cancer 1	BRCA1
•	Hereditary breast/ovarian cancer 2	BRCA2
Sarcoma	Hereditary retinoblastoma	RB1
	Li-Fraumeni syndrome	TP53
	Neurofibromatosis 1	NF1
Skin	Hereditary melanoma 1	CDKN2
	Hereditary melanoma 2	CDK4
	Basal cell naevus (Gorlin) syndrome	PTCH
Stomach	Hereditary non-polyposis colon cancer (HNPCC) 1	hMSH2
	Hereditary non-polyposis colon cancer (HNPCC) 2	hMLH1
	Hereditary non-polyposis colon cancer (HNPCC) 3	hPMS1
	Hereditary non-polyposis colon cancer (HNPCC) 4	hPMS2

The polynucleotides of the invention can be especially useful to monitor patients having any of the above syndromes to detect potentially malignant events at a molecular level before they are detectable at a gross morphological level. As can be seen from the table, a number of genes are involved in multiple forms of cancer. Thus, a polynucleotide of the invention identified as important for metastatic colon cancer can also have clinical implications for a patient diagnosed with stomach cancer or endometrial cancer.

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Lung Cancer. Lung cancer is one of the most common cancers in the United States, accounting for about 15 percent of all cancer cases, or 170,000 new cases each year. At this time, over half of the lung cancer cases in the United States are in men, but the number found in women is increasing and will soon equal that in men. Today more women die of lung cancer than of breast cancer. Lung cancer is especially difficult to diagnose and treat because of the large size of the lungs, which allows cancer to develop for years undetected. In fact, lung cancer can spread outside the lungs without causing any symptoms. Adding to the confusion, the most common symptom of lung cancer, a persistent cough, can often be mistaken for a cold or bronchitis.

Although there are more than a dozen different kinds of lung cancer, the two main types of lung cancer are small cell and nonsmall cell, which encompass about 90% of all lung cancer cases. Small cell carcinoma (also called oat cell carcinoma), which usually starts in one of the larger bronchial tubes, grows fairly rapidly, and is likely to be large by the time of diagnosis. Nonsmall cell lung cancer (NSCLC) is made up of three general subtypes of lung cancer. Epidermoid carcinoma (also called squamous cell carcinoma) usually starts in one of the larger bronchial tubes and grows relatively slowly. The size of these tumors can range from very small to quite large. Adenocarcinoma starts growing near the outside surface of the lung and can vary in both size and growth rate. Some slowly growing adenocarcinomas are described as alveolar cell cancer. Large cell carcinoma starts near the surface of the lung, grows rapidly, and the growth is usually fairly large when diagnosed. Other less common forms of lung cancer are carcinoid, cylindroma, mucoepidermoid, and malignant mesothelioma.

Currently, CT scans, MRIs, X-rays, sputum cytology, and biopsies are used to diagnose nonsmall cell lung cancer. The form and cellular origin of the lung cancer is diagnosed primarily through biopsy from either a surgical biopsy or a needle aspiration of lung tissue, and usually the biopsy is prompted from an abnormality identified on an X-ray. In some cases, sputum cytology can reveal lung cancers in patients with normal X-rays or can determine the type of lung cancer, but because it cannot pinpoint the tumor's location, a positive sputum cytology test is usually followed by further tests. Since these tests are based in large part on gross morphology of the tissue, the diagnosis of a particular kind of tumor is largely subjective, and the diagnosis can vary significantly between clinicians.

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The polynucleotides of the invention can be used to distinguish types of lung cancer as well as identifying traits specific to a certain patient's cancer. For example, if the patient's biopsy expresses a polynucleotide that is associated with a low metastatic potential, it may justify leaving a larger portion of the patient's lung in surgery to remove the lesion. Alternatively, a smaller lesion with expression of a polynucleotide that is associated with high metastatic potential may justify a more radical removal of lung tissue and/or the surrounding lymph nodes, even if no metastasis can be identified through pathological examination.

Similarly, the expression of polynucleotides of the invention can be used in the diagnosis, prognosis and management of colorectal cancer. The differential expression of a polynucleotide in hyperplasia can be used as a diagnostic marker for metastatic lung cancer. The polynucleotides of the invention that would be especially useful for this purpose are those that exhibit differential expression between high metastatic versus low metastatic lung cancer, *i.e.* SEQ ID NOS: 174, 254, 466, 571, 574, 590, 922, 1355, 1422, 2007, 2038, 2245, 10, 54, 65, 171, 203, 252, 253, 285, 419, 420, 491, 525, 526, 552, 693, 700, 726, 742, 746, 861, 990, 1088, 1288, 1417, 1444, 1454, 1570, 1597, 1979, 2024, 2034, and 2126. Detection of malignant lung cancer with a higher metastatic potential can be determined using expression levels of any of these sequences alone or in combination with the levels of expression of other known genes.

Breast Cancer. The National Cancer Institute (NCI) estimates that about 1 in 8 women in the United States will develop breast cancer during her lifetime. Clinical breast examination and mammography are recommended as combined modalities for breast cancer screening, and the nature of the cancer will often depend upon the location of the tumor and the cell type from which the tumor is derived. The majority of breast cancers are adenocarcinomas subtypes, which can be summarized as follows:

Ductal carcinoma in situ (DCIS): Ductal carcinoma in situ is the most common type of noninvasive breast cancer. In DCIS, the malignant cells have not metastasized through the walls of the ducts into the fatty tissue of the breast. Comedocarcinoma is a type of DCIS that is more likely than other types of DCIS to come back in the same area after lumpectomy. It is more closely linked to eventual development of invasive ductal carcinoma than other forms of DCIS.

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Infiltrating (or invasive) ductal carcinoma (IDC): this type of cancer has metastasized through the wall of the duct and invaded the fatty tissue of the breast. At this point, it has the potential to use the lymphatic system and bloodstream for metastasis to more distant parts of the body. Infiltrating ductal carcinoma accounts for about 80% of breast cancers.

Lobular carcinoma in situ (LCIS): While not a true cancer, LCIS (also called lobular neoplasia) is sometimes classified as a type of noninvasive breast cancer. It does not penetrate through the wall of the lobules. Although it does not itself usually become an invasive cancer, women with this condition have a higher risk of developing an invasive breast cancer in the same breast, or in the opposite breast.

Infiltrating (or invasive) lobular carcinoma (ILC): ILC is similar to IDC, in that it has the potential metastasize elsewhere in the body. About 10% to 15% of invasive breast cancers are invasive lobular carcinomas. ILC can be more difficult to detect by mammogram than IDC.

Inflammatory breast cancer: This rare type of invasive breast cancer accounts for about 1% of all breast cancers and is extremely aggressive. Multiple skin symptoms associated with this cancer are caused by cancer cells blocking lymph vessels or channels in the skin over the breast.

Medullary carcinoma: This special type of infiltrating breast cancer has a relatively well defined, distinct boundary between tumor tissue and normal tissue. It accounts for about 5% of breast cancers. The prognosis for this kind of breast cancer is better than for other types of invasive breast cancer.

Mucinous carcinoma: This rare type of invasive breast cancer originates from mucus-producing cells. The prognosis for mucinous carcinoma is better than for the more common types of invasive breast cancer.

Paget's disease of the nipple: This type of breast cancer starts in the ducts and spreads to the skin of the nipple and the areola. It is a rare type of breast cancer, occurring in only 1% of all cases. Paget's disease can be associated with in situ carcinoma, or with infiltrating breast carcinoma. If no lump can be felt in the breast tissue, and the biopsy shows DCIS but no invasive cancer, the prognosis is excellent.

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Phyllodes tumor: This very rare type of breast tumor forms from the stroma of the breast, in contrast to carcinomas which develop in the ducts or lobules. Phyllodes (also spelled phylloides) tumors are usually benign, but are malignant on rare occasions. Nevertheless, malignant phyllodes tumors are very rare and less than 10 women per year in the US die of this disease. Benign phyllodes tumors are successfully treated by removing the mass and a narrow margin of normal breast tissue.

Tubular carcinoma: Accounting for about 2% of all breast cancers, tubular carcinomas are a special type of infiltrating breast carcinoma. They have a better prognosis than usual infiltrating ductal or lobular carcinomas.

High-quality mammography combined with clinical breast exam remains the only screening method clearly tied to reduction in breast cancer mortality. Lower dose x-rays, digitized computer rather than film images, and the use of computer programs to assist diagnosis, are almost ready for widespread dissemination. Other technologies also are being developed, including magnetic resonance imaging and ultrasound. In addition, a very low radiation exposure technique, positron emission tomography has the potential for detecting early breast cancer.

It is also possible to differentiate between non-cancerous breast tissue and malignant breast tissue by analyzing differential gene expression between tissues. In addition, there may be several possible alterations that lead to the various possible types of breast cancer. The different types of breast tumors (e.g., invasive vs. non-invasive, ductal vs. axillary lymph node) can be differentiable from one another by the identification of the differences in genes expressed by different types of breast tumor tissues (Porter-Jordan et al., Hematol Oncol Clin North Am (1994) 8:73). Breast cancer can thus be generally diagnosed by detection of expression of a gene or genes associated with breast tumors. Where enough information is available about the differential gene expression between various types of breast tumor tissues, the specific type of breast tumor can also be diagnosed.

For example, increased estrogen receptor (ER) expression in normal breast epithileum, while not itself indicative of malignant tissue, is a known risk marker for development of breast cancer. Khan SA et al., Cancer Res (1994) 54:993. Malignant breast cancer is often divided into two groups, ER-positive and ER-negative, based on the

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estrogen receptor status of the tissue. The ER status represents different survival length and response to hormone therapy, and is thought to represent either: 1) an indicator of different stages of the disease, or 2) an indicator that allows differentiation between two similar but distinct diseases. K. Zhu *et al.*, *Med. Hypoth.* (1997) 49:69. A number of other genes are known to vary expression between either different stages of cancer or different types of similar breast cancer.

Similarly, the expression of polynucleotides of the invention can be used in the diagnosis and management of breast cancer. The differential expression of a polynucleotide in human breast tumor tissue can be used as a diagnostic marker for human breast cancer. The polynucleotides of the invention that would be especially useful for this purpose are those that exhibit differential expression between breast cancer tissue with a high metastatic potential and a low metastatic potential, i.e. SEQ ID NOS:15, 36, 44, 89, 172, 203, 261, 419, 420, 503, 552, 564, 570, 590, 693, 707, 711, 726, 746, 756, 990, 1122, 1142, 1286, 1289, 1435, 1860, 1933, 1934, 1979, 1980, 2007, 2023, 2409, 2486, 45, 146, 154, 159, 165, 174, 183, 364, 366, 387, 496, 510, 512, 529, 560, 606, 644, 646, 754, 875, 902, 921, 942, 1095, 1104, 1131, 1170, 1184, 1205, 1354, 1387, 1535, 1751, 1764, 1777, 1795, 1869, 1882, 1890, 1915, 2040, 2059, 2223, 2245, 2300, 2325, 2462, 2488, 2492; Detection of breast cancer can be determined using expression levels of any of these sequences alone or in combination. Determination of the aggressive nature and/or the metastatic potential of a breast cancer can also be determined by comparing levels of one or more polynucleotides of the invention and comparing levels of another sequence known to vary in cancerous tissue, e.g. ER expression. In addition, development of breast cancer can be detected by examining the ratio of SEQ ID NO: to the levels of steroid hormones (e.g., testosterone or estrogen) or to other hormones (e.g., growth hormone, insulin). Thus expression of specific marker polynucleotides can be used to discriminate between normal and cancerous breast tissue, to discriminate between breast cancers with different cells of origin, to discriminate between breast cancers with different potential metastatic rates, etc.

Diagnosis of breast cancer can also involve comparing the expression of a polynucleotide of the invention with the expression of other sequences in non-malignant breast tissue samples in comparison to one or more forms of the diseased tissue. A comparison of expression of one or more polynucleotides of the invention between the

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samples provides information on relative levels of these polynucleotides as well as the ratio of these polynucleotides to the expression of other sequences in the tissue of interest compared to normal.

This risk of breast cancer is elevated significantly by the presence of an inherited risk for breast cancer, such as a mutation in BRCA-1 or BRCA-2. New diagnostic tools are being developed to address the needs of higher risk patients to complement mammography and physical examinations for early detection of breast cancer, particularly among younger women. The presence of antigen or expression markers in nipple aspirate fluid (NAF) samples collected from one or both breasts can be useful for useful for risk assessment or early cancer detection. Breast cytology and biomarkers obtained by random fine needle aspiration have been used to identify hyperplasia with atypia and overexpression of p53 and EGFR. The polynucleotides of the invention can be used in multivariate analysis with expression studies with genes such as p53 and EGFR as risk predictors and as surrogate endpoint biomarkers for breast cancer.

As well as being used for diagnosis and risk assessment, the expression of certain genes can also correlated to prognosis of a disease state. The expression of particular gene have been used as prognostic indicators for breast cancer including increased expression of c-erbB-2, pS2, ER, progesterone receptor, epidermal growth factor receptor (EGFR), neu, myc, bcl-2, int2, cytosolic tyrosine kinase, cyclin E, prad-1, hst, uPA, PAI-1, PAI-2, cathepsin D, as well as the presence of a number of cancer-specific antigens, e.g. CEA, CA M26, CA M29 and CA 15.3. Davis, Br. J. Biomed Sci. (1996) 53:157. Poor prognosis has also been linked to a decrease in expression of certain genes, such as p53, Rb, nm23. The expression of the polynucleotides of the invention can be of prognostic value for determining the metastatic potential of a malignant breast cancer, as this molecules are differentially expressed between high and low metastatic potential tissues tumors. The levels of these polynucleotides in patients with malignant breast cancer can compared to normal tissue, malignant tissue with a known high potential metastatic level, and malignant tissue with a known lower level of metastatic potential to provide a prognosis for a particular patient. Such a prognosis is predictive of the extent and nature of the cancer. The determined prognosis is useful in determining the prognosis of a patient with breast cancer, both for initial treatment of the disease and for longer-term monitoring of the same

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patient. If samples are taken from the same individual over a period of time, differences in polynucleotide expression that are specific to that patient can be identified and closely watched.

Colon Cancer. Colorectal cancer is one of the most common neoplasms in humans and perhaps the most frequent form of hereditary neoplasia. Prevention and early detection are key factors in controlling and curing colorectal cancer. Indeed, colorectal cancer is the second most preventable cancer, after lung cancer. Colorectal cancer begins as polyps, which are small, benign growths of cells that form on the inner lining of the colon. Over a period of several years, some of these polyps accumulate additional mutations and become cancerous. About 20 percent of all cases of colon cancer are thought to be related to heredity. Currently, multiple familial colorectal cancer disorders have been identified, which are summarized as follows:

Familial adenomatous polyposis (FAP): This condition results in a person having hundreds or even thousands of polyps in the colon and rectum that usually first appear during the teenage years. Cancer nearly always develops in one or more of these polyps between the ages of 30 and 50.

Gardner's syndrome: Like FAP, Gardner's syndrome results in polyps and colorectal cancers that develop at a young age. It can also cause benign tumors of the skin, soft connective tissue and bones.

Hereditary nonpolyposis colon cancer (HNPCC): People with this condition tend to develop colorectal cancer at a young age, without first having many polyps. HNPCC has an autosomal dominant pattern of inheritance with variable but high penetrance estimated to be about 90%. HNPCC underlies 0.5%-10% of all cases of colorectal cancer. An understanding of the mechanisms behind the development of HNPCC is emerging, and genetic presymptomatic testing, now being conducted in research settings, soon will be available on a widespread basis for individuals identified at risk for this disease.

Familial colorectal cancer in Ashkenazi Jews: Recent research has found an inherited tendency to developing colorectal cancer among some Jews of Eastern European descent. Like people with FAP, Gardner's syndrome, and HNPCC, their increased risk is due to an inherited mutation present in about 6% of American Jews.

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Several tests are currently used to screen for colorectal cancer, including digital rectal examination, fecal occult blood test, sigmoidoscopy, colonoscopy, virtual colonoscopy and MRI. Each of these tests identifies potential colorectal cancer lesions, or a risk of development of these lesions, at a fairly gross morphological level.

The sequential alteration of a number of genes is associated with malignant adenocarcinoma, including the genes DCC, p53, ras, and FAP. For a review, see *e.g.*Fearon ER, *et al.*, *Cell* (1990) *61(5)*:759; Hamilton SR *et al.*, *Cancer* (1993) 72:957;
Bodmer W, *et al.*, *Nat Genet.* (1994) *4(3)*:217; Fearon ER, *Ann N Y Acad Sci.* (1995) 768:101. Molecular genetic alterations are thus promising as potential diagnostic and prognostic indicators in colorectal carcinoma and molecular genetics of colorectal carcinoma since it is possible to differentiate between different types of colorectal neoplasias using molecular markers. Colorectal cancer can thus be generally diagnosed by detection of expression of a gene or genes associated with colorectal tumors.

Similarly, the expression of polynucleotides of the invention can be used in the diagnosis, prognosis and management of colorectal cancer. The differential expression of a polynucleotide in hyperplasia can be used as a diagnostic marker for colon cancer. The polynucleotides of the invention that would be especially useful for this purpose are those that exhibit differential expression between malignant metastatic colon cancer and normal patient tissue, *i.e.* SEQ ID NOS:228, 280, 355, 491, 603, 680, 752, 753, 1241, 1264, 1401, 1442, 1514, 1851, 1915, 2024, 2066, 33, 250, 282, 370, 387, 443, 460, 545, 560, 703, 704, 1095, 1104, 1205, 1354, 1387, 1734, 1742, 1954, 2262, 2325, 1899, 252, 253, 491, 581, 693, 726, 746, 1780, 1899, 65, 252, 253, 581, 693, 716, 726, 746, 1780, 1899, and 1780. Detection of malignant colon cancer can be determined using expression levels of any of these sequences alone or in combination with the levels of expression.

Determination of the aggressive nature and/or the metastatic potential of a colon cancer can also be determined by comparing levels of one or more polynucleotides of the invention and comparing total levels of another sequence known to vary in cancerous tissue, e.g. p53 expression. In addition, development of colon cancer can be detected by examining the ratio of any of the polynucleotides of the invention to the levels of oncogenes (e.g. ras) or tumor suppressor genes (e.g. FAP or p53). Thus expression of specific marker polynucleotides can be used to discriminate between normal and cancerous

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breast tissue, to discriminate between breast cancers with different cells of origin, to discriminate between breast cancers with different potential metastatic rates, etc.

G. <u>Use of Polynucleotides to Screen for Peptide Analogs and Antagonists</u>

Polypeptides encoded by the instant polynucleotides and corresponding full length genes can be used to screen peptide libraries to identify binding partners, such as receptors, from among the encoded polypeptides.

A library of peptides can be synthesized following the methods disclosed in U.S. Pat. No. 5,010,175 ('175), and in WO 91/17823. As described below in brief, one prepares a mixture of peptides, which is then screened to identify the peptides exhibiting the desired signal transduction and receptor binding activity. In the '175 method, a suitable peptide synthesis support (e.g., a resin) is coupled to a mixture of appropriately protected, activated amino acids. The concentration of each amino acid in the reaction mixture is balanced or adjusted in inverse proportion to its coupling reaction rate so that the product is an equimolar mixture of amino acids coupled to the starting resin. The bound amino acids are then deprotected, and reacted with another balanced amino acid mixture to form an equimolar mixture of all possible dipeptides. This process is repeated until a mixture of peptides of the desired length (e.g., hexamers) is formed. Note that one need not include all amino acids in each step: one can include only one or two amino acids in some steps (e.g., where it is known that a particular amino acid is essential in a given position), thus reducing the complexity of the mixture. After the synthesis of the peptide library is completed, the mixture of peptides is screened for binding to the selected polypeptide. The peptides are then tested for their ability to inhibit or enhance activity. Peptides exhibiting the desired activity are then isolated and sequenced.

The method described in WO 91/17823 is similar. However, instead of reacting the synthesis resin with a mixture of activated amino acids, the resin is divided into twenty equal portions (or into a number of portions corresponding to the number of different amino acids to be added in that step), and each amino acid is coupled individually to its portion of resin. The resin portions are then combined, mixed, and again divided into a number of equal portions for reaction with the second amino acid. In this manner, each reaction can be easily driven to completion. Additionally, one can maintain separate "subpools" by treating portions in parallel, rather than combining all resins at each step.

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This simplifies the process of determining which peptides are responsible for any observed receptor binding or signal transduction activity.

In such cases, the subpools containing, e.g., 1-2,000 candidates each are exposed to one or more polypeptides of the invention. Each subpool that produces a positive result is then resynthesized as a group of smaller subpools (sub-subpools) containing, e.g., 20-100 candidates, and reassayed. Positive sub-subpools can be resynthesized as individual compounds, and assayed finally to determine the peptides that exhibit a high binding constant. These peptides can be tested for their ability to inhibit or enhance the native activity. The methods described in WO 91/7823 and U.S. Patent No. 5,194,392 (herein incorporated by reference) enable the preparation of such pools and subpools by automated techniques in parallel, such that all synthesis and resynthesis can be performed in a matter of days.

Peptide agonists or antagonists are screened using any available method, such as signal transduction, antibody binding, receptor binding, mitogenic assays, chemotaxis assays, etc. The methods described herein are presently preferred. The assay conditions ideally should resemble the conditions under which the native activity is exhibited *in vivo*, that is, under physiologic pH, temperature, and ionic strength. Suitable agonists or antagonists will exhibit strong inhibition or enhancement of the native activity at concentrations that do not cause toxic side effects in the subject. Agonists or antagonists that compete for binding to the native polypeptide can require concentrations equal to or greater than the native concentration, while inhibitors capable of binding irreversibly to the polypeptide can be added in concentrations on the order of the native concentration.

The end results of such screening and experimentation will be at least one novel polypeptide binding partner, such as a receptor, encoded by a gene or a cDNA corresponding to a polynucleotide of the invention, and at least one peptide agonist or antagonist of the novel binding partner. Such agonists and antagonists can be used to modulate, enhance, or inhibit receptor function in cells to which the receptor is native, or in cells that possess the receptor as a result of genetic engineering. Further, if the novel receptor shares biologically important characteristics with a known receptor, information about agonist/antagonist binding can facilitate development of improved agonists/antagonists of the known receptor.

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H. Pharmaceutical Compositions and Therapeutic Uses

Pharmaceutical compositions can comprise polypeptides, antibodies, or polynucleotides of the claimed invention. The pharmaceutical compositions will comprise a therapeutically effective amount of either polypeptides, antibodies, or polynucleotides of the claimed invention.

The term "therapeutically effective amount" as used herein refers to an amount of a therapeutic agent to treat, ameliorate, or prevent a desired disease or condition, or to exhibit a detectable therapeutic or preventative effect. The effect can be detected by, for example, chemical markers or antigen levels. Therapeutic effects also include reduction in physical symptoms, such as decreased body temperature. The precise effective amount for a subject will depend upon the subject's size and health, the nature and extent of the condition, and the therapeutics or combination of therapeutics selected for administration. Thus, it is not useful to specify an exact effective amount in advance. However, the effective amount for a given situation is determined by routine experimentation and is within the judgment of the clinician. For purposes of the present invention, an effective dose will generally be from about 0.01 mg/kg to 50 mg/kg or 0.05 mg/kg to about 10 mg/kg of the DNA constructs in the individual to which it is administered.

A pharmaceutical composition can also contain a pharmaceutically acceptable carrier. The term "pharmaceutically acceptable carrier" refers to a carrier for administration of a therapeutic agent, such as antibodies or a polypeptide, genes, and other therapeutic agents. The term refers to any pharmaceutical carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition, and which can be administered without undue toxicity. Suitable carriers can be large, slowly metabolized macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, and inactive virus particles. Such carriers are well known to those of ordinary skill in the art.

Pharmaceutically acceptable salts can be used therein, for example, mineral acid salts such as hydrochlorides, hydrobromides, phosphates, sulfates, and the like; and the salts of organic acids such as acetates, propionates, malonates, benzoates, and the like. A thorough discussion of pharmaceutically acceptable excipients is available in *Remington's Pharmaceutical Sciences* (Mack Pub. Co., N.J. 1991).

Pharmaceutically acceptable carriers in therapeutic compositions can include liquids such as water, saline, glycerol and ethanol. Auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, can also be present in such vehicles. Typically, the therapeutic compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection can also be prepared. Liposomes are included within the definition of a pharmaceutically acceptable carrier.

Delivery Methods. Once formulated, the compositions of the invention can be (1) administered directly to the subject (e.g., as polynucleotide or polypeptides); (2) delivered ex vivo, to cells derived from the subject (e.g., as in ex vivo gene therapy); or (3) delivered in vitro for expression of recombinant proteins (e.g., polynucleotides). Direct delivery of the compositions will generally be accomplished by injection, either subcutaneously, intraperitoneally, intravenously or intramuscularly, or delivered to the interstitial space of a tissue. The compositions can also be administered into a tumor or lesion. Other modes of administration include oral and pulmonary administration, suppositories, and transdermal applications, needles, and gene guns or hyposprays. Dosage treatment can be a single dose schedule or a multiple dose schedule.

Methods for the ex vivo delivery and reimplantation of transformed cells into a subject are known in the art and described in *e.g.*, International Publication No. WO 93/14778. Examples of cells useful in ex vivo applications include, for example, stem cells, particularly hematopoetic, lymph cells, macrophages, dendritic cells, or tumor cells. Generally, delivery of nucleic acids for both ex vivo and in vitro applications can be accomplished by, for example, dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei, all well known in the art.

Once a gene corresponding to a polynucleotide of the invention has been found to correlate with a proliferative disorder, such as neoplasia, dysplasia, and hyperplasia, the disorder can be amenable to treatment by administration of a therapeutic agent based on the provided polynucleotide or corresponding polypeptide.

Preparation of antisense polynucleotides is discussed above. Neoplasias that are treated with the antisense composition include, but are not limited to, cervical cancers, melanomas, colorectal adenocarcinomas, Wilms' tumor, retinoblastoma, sarcomas, myosarcomas, lung carcinomas, leukemias, such as chronic myelogenous leukemia, promyelocytic leukemia, monocytic leukemia, and myeloid leukemia, and lymphomas, such as histiocytic lymphoma. Proliferative disorders that are treated with the therapeutic composition include disorders such as anhydric hereditary ectodermal dysplasia, congenital alveolar dysplasia, epithelial dysplasia of the cervix, fibrous dysplasia of bone, and mammary dysplasia. Hyperplasias, for example, endometrial, adrenal, breast, prostate, or thyroid hyperplasias or pseudoepitheliomatous hyperplasia of the skin, are treated with antisense therapeutic compositions based upon a polynucleotide of the invention. Even in disorders in which mutations in the corresponding gene are not implicated, downregulation or inhibition of expression of a gene corresponding to a polynucleotide of the invention can have therapeutic application. For example, decreasing gene expression can help to suppress tumors in which enhanced expression of the gene is implicated.

Both the dose of the antisense composition and the means of administration are determined based on the specific qualities of the therapeutic composition, the condition, age, and weight of the patient, the progression of the disease, and other relevant factors. Administration of the therapeutic antisense agents of the invention includes local or systemic administration, including injection, oral administration, particle gun or catheterized administration, and topical administration. Preferably, the therapeutic antisense composition contains an expression construct comprising a promoter and a polynucleotide segment of at least 12, 22, 25, 30, or 35 contiguous nucleotides of the antisense strand of a polynucleotide disclosed herein. Within the expression construct, the polynucleotide segment is located downstream from the promoter, and transcription of the polynucleotide segment initiates at the promoter.

Various methods are used to administer the therapeutic composition directly to a specific site in the body. For example, a small metastatic lesion is located and the therapeutic composition injected several times in several different locations within the body of tumor. Alternatively, arteries which serve a tumor are identified, and the therapeutic composition injected into such an artery, in order to deliver the composition directly into

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the tumor. A tumor that has a necrotic center is aspirated and the composition injected directly into the now empty center of the tumor. The antisense composition is directly administered to the surface of the tumor, for example, by topical application of the composition. X-ray imaging is used to assist in certain of the above delivery methods.

Receptor-mediated targeted delivery of therapeutic compositions containing an antisense polynucleotide, subgenomic polynucleotides, or antibodies to specific tissues is also used. Receptor-mediated DNA delivery techniques are described in, for example, Findeis et al., Trends Biotechnol. (1993) 11:202; Chiou et al., Gene Therapeutics: Methods And Applications Of Direct Gene Transfer (J.A. Wolff, ed.) (1994); Wu et al., J. Biol. Chem. (1988) 263:621; Wu et al., J. Biol. Chem. (1994) 269:542; Zenke et al., Proc. Natl. Acad. Sci. (USA) (1990) 87:3655; Wu et al., J. Biol. Chem. (1991) 266:338. Preferably, receptor-mediated targeted delivery of therapeutic compositions containing antibodies of the invention is used to deliver the antibodies to specific tissue.

Therapeutic compositions containing antisense subgenomic polynucleotides are administered in a range of about 100 ng to about 200 mg of DNA for local administration in a gene therapy protocol. Concentration ranges of about 500 ng to about 50 mg, about 1 µg to about 2 mg, about 5 µg to about 500 µg, and about 20 µg to about 100 µg of DNA can also be used during a gene therapy protocol. Factors such as method of action and efficacy of transformation and expression are considerations which will affect the dosage required for ultimate efficacy of the antisense subgenomic polynucleotides. Where greater expression is desired over a larger area of tissue, larger amounts of antisense subgenomic polynucleotides or the same amounts readministered in a successive protocol of administrations, or several administrations to different adjacent or close tissue portions of, for example, a tumor site, may be required to effect a positive therapeutic outcome. In all cases, routine experimentation in clinical trials will determine specific ranges for optimal therapeutic effect. A more complete description of gene therapy vectors, especially retroviral vectors, is contained in U.S. Serial No. 08/869,309, which is expressly incorporated herein, and in section G below.

For polynucleotide-related genes encoding polypeptides or proteins with antiinflammatory activity, suitable use, doses, and administration are described in U.S. Patent No. 5,654,173. Therapeutic agents also include antibodies to proteins and polypeptides

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encoded by the polynucleotides of the invention and related genes, as described in U.S. Patent No. 5,654,173.

I. Gene Therapy

The therapeutic polynucleotides and polypeptides of the present invention can be utilized in gene delivery vehicles. The gene delivery vehicle can be of viral or non-viral origin (see generally, Jolly, Cancer Gene Therapy (1994) 1:51; Kimura, Human Gene Therapy (1994) 5:845; Connelly, Human Gene Therapy (1995) 1:185; and Kaplitt, Nature Genetics (1994) 6:148). Gene therapy vehicles for delivery of constructs including a coding sequence of a therapeutic of the invention can be administered either locally or systemically. These constructs can utilize viral or non-viral vector approaches. Expression of such coding sequences can be induced using endogenous mammalian or heterologous promoters. Expression of the coding sequence can be either constitutive or regulated.

The present invention can employ recombinant retroviruses which are constructed to carry or express a selected nucleic acid molecule of interest. Retrovirus vectors that can be employed include those described in EP 0 415 731; WO 90/07936; WO 94/03622; WO 93/25698; WO 93/25234; U.S. Patent No. 5, 219,740; WO 93/11230; WO 93/10218; Vile and Hart, Cancer Res. (1993) 53:3860; Vile et al., Cancer Res. (1993) 53:962; Ram et al., Cancer Res. (1993) 53:83; Takamiya et al., J. Neurosci. Res. (1992) 33:493; Baba et al., J. Neurosurg. (1993) 79:729; U.S. Patent No. 4,777,127; GB Patent No. 2,200,651; and EP 0 345 242. Preferred recombinant retroviruses include those described in WO 91/02805.

Packaging cell lines suitable for use with the above-described retroviral vector constructs can be readily prepared (see, e.g., WO 95/30763 and WO 92/05266), and used to create producer cell lines (also termed vector cell lines) for the production of recombinant vector particles. Within particularly preferred embodiments of the invention, packaging cell lines are made from human (such as HT1080 cells) or mink parent cell lines, thereby allowing production of recombinant retroviruses that can survive inactivation in human serum.

The present invention also employs alphavirus-based vectors that can function as gene delivery vehicles. Such vectors can be constructed from a wide variety of alphaviruses, including, for example, Sindbis virus vectors, Semliki forest virus (ATCC VR-67; ATCC VR-1247), Ross River virus (ATCC VR-373; ATCC VR-1246) and

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Venezuelan equine encephalitis virus (ATCC VR-923; ATCC VR-1250; ATCC VR 1249; ATCC VR-532). Representative examples of such vector systems include those described in U.S. Patent Nos. 5,091,309; 5,217,879; and 5,185,440; WO 92/10578; WO 94/21792; WO 95/27069; WO 95/27044; and WO 95/07994. Gene delivery vehicles of the present invention can also employ parvovirus such as adeno-associated virus (AAV) vectors. Representative examples include the AAV vectors disclosed by Srivastava in WO 93/09239, Samulski et al., *J. Virol.* (1989) 63:3822; Mendelson et al., *Virol.* (1988) 166:154; and Flotte et al., PNAS (1993) 90:10613.

Representative examples of adenoviral vectors include those described by Berkner,

Biotechniques (1988) 6:616; Rosenfeld et al., Science (1991) 252:431; WO 93/19191;

Kolls et al., PNAS (1994) 91:215; Kass-Eisler et al., PNAS (1993) 90:11498; Guzman et al., Circulation (1993) 88:2838; Guzman et al., Cir. Res. (1993) 73:1202; Zabner et al., Cell (1993) 75:207; Li et al., Hum. Gene Ther. (1993) 4:403; Cailaud et al., Eur. J. Neurosci. (1993) 5:1287; Vincent et al., Nat. Genet. (1993) 5:130; Jaffe et al., Nat. Genet. (1992) 1:372; and Levrero et al., Gene (1991) 101:195. Exemplary adenoviral gene therapy vectors employable in this invention also include those described in WO 94/12649, WO 93/03769; WO 93/19191; WO 94/28938; WO 95/11984 and WO 95/00655. Administration of DNA linked to killed adenovirus as described in Curiel, Hum. Gene Ther. (1992) 3:147 can be employed.

Other gene delivery vehicles and methods can be employed, including polycationic condensed DNA linked or unlinked to killed adenovirus alone, for example Curiel, *Hum. Gene Ther.* (1992) 3:147; ligand linked DNA, for example see Wu, *J. Biol. Chem.* (1989) 264:16985; eukaryotic cell delivery vehicles cells, for example see U.S. Pat. No. 5,814,482; WO 95/07994; WO 96/17072; WO 95/30763; and WO 97/42338; deposition of photopolymerized hydrogel materials; hand-held gene transfer particle gun, as described in U.S. Patent No. 5,149,655; ionizing radiation as described in U.S. Patent No. 5,206,152 and in WO92/11033; nucleic charge neutralization or fusion with cell membranes. Additional approaches are described in Philip, *Mol. Cell Biol.* (1994) 14:2411, and in Woffendin, *Proc. Natl. Acad. Sci.* (1994) 91:1581.

Naked DNA can also be employed. Exemplary naked DNA introduction methods are described in WO 90/11092 and U.S. Patent No. 5,580,859. Liposomes that can act as

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gene delivery vehicles are described in U.S. Patent No. 5,422,120; WO 95/13796; WO 94/23697; WO 91/14445; and EP 0524968.

Further non-viral delivery suitable for use includes mechanical delivery systems such as the approach described in Woffendin *et al.*, *Proc. Natl. Acad. Sci. USA* (1994) 91(24):11581. Moreover, the coding sequence and the product of expression of such can be delivered through deposition of photopolymerized hydrogel materials. Other conventional methods for gene delivery that can be used for delivery of the coding sequence include, for example, use of hand-held gene transfer particle gun, as described in U.S. Patent No. 5,149,655; use of ionizing radiation for activating transferred gene, as described in U.S. Patent No. 5,206,152 and WO 92/11033.

The present invention will now be illustrated by reference to the following examples which set forth particularly advantageous embodiments. However, it should be noted that these embodiments are illustrative and are not to be construed as restricting the invention in any way.

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EXAMPLES

Example 1: Source of Biological Materials and Overview of Novel Polynucleotides

Expressed by the Biological Materials

Human colon cancer cell line Km12L4-A (Morika, W. A. K. et al., *Cancer Research* (1988) 48:6863) was used to construct a cDNA library from mRNA isolated from the cells. As described in the above overview, a total of 4,693 sequences expressed by the Km12L4-A cell line were isolated and analyzed; most sequences were about 275-300 nucleotides in length. The KM12L4-A cell line is derived from the KM12C cell line. The KM12C cell line, which is poorly metastatic (low metastatic) was established in culture from a Dukes' stage B₂ surgical specimen (Morikawa *et al. Cancer Res.* (1988) 48:6863). The KML4-A is a highly metastatic subline derived from KM12C (Yeatman *et al. Nucl. Acids. Res.* (1995) 23:4007; Bao-Ling *et al. Proc. Annu. Meet. Am. Assoc. Cancer. Res.* (1995) 21:3269). The KM12C and KM12C-derived cell lines (*e.g.*, KM12L4, KM12L4-A, *etc.*) are well-recognized in the art as a model cell line for the study of colon cancer (see, *e.g.*, Moriakawa *et al.*, *supra*; Radinsky *et al. Clin. Cancer Res.* (1995) 1:19; Yeatman *et al.* (1995) *supra*; Yeatman *et al. Clin. Exp. Metastasis* (1996) 14:246).

The sequences were first masked to eliminate low complexity sequences using the XBLAST masking program (Claverie "Effective Large-Scale Sequence Similarity Searches," In: Computer Methods for Macromolecular Sequence Analysis, Doolittle, ed., Meth. Enzymol. 266:212-227 Academic Press, NY, NY (1996); see particularly Claverie, in "Automated DNA Sequencing and Analysis Techniques" Adams et al., eds., Chap. 36, p. 267 Academic Press, San Diego, 1994 and Claverie et al. Comput. Chem. (1993) 17:191). Generally, masking does not influence the final search results, except to eliminate sequences of relative little interest due to their low complexity, and to eliminate multiple "hits" based on similarity to repetitive regions common to multiple sequences, e.g., Alu repeats. Masking resulted in the elimination of 43 sequences. The remaining sequences were then used in a BLASTN vs. Genbank search with search parameters of greater than 70% overlap, 99% identity, and a p value of less than 1 x 10⁻⁴⁰, which search resulted in the discarding of 1,432 sequences. Sequences from this search also were discarded if the inclusive parameters were met, but the sequence was ribosomal or vector-derived.

The resulting sequences from the previous search were classified into three groups (1, 2 and 3 below) and searched in a BLASTX vs. NRP (non-redundant proteins) database

search: (1) unknown (no hits in the Genbank search), (2) weak similarity (greater than 45% identity and p value of less than 1×10^{-5}), and (3) high similarity (greater than 60% overlap, greater than 80% identity, and p value less than 1×10^{-5}). This search resulted in discard of 98 sequences as having greater than 70% overlap, greater than 99% identity, and p value of less than 1×10^{-40} .

The remaining sequences were classified as unknown (no hits), weak similarity, and high similarity (parameters as above). Two searches were performed on these sequences. First, a BLAST vs. EST database search resulted in discard of 1771 sequences (sequences with greater than 99% overlap, greater than 99% similarity and a p value of less than 1 x 10^{-40} ; sequences with a p value of less than 1 x 10^{-65} when compared to a database sequence of human origin were also excluded). Second, a BLASTN vs. Patent GeneSeq database resulted in discard of 15 sequences (greater than 99% identity; p value less than 1 x 10^{-40} ; greater than 99% overlap).

The remaining sequences were subjected to screening using other rules and redundancies in the dataset. Sequences with a p value of less than 1 x 10 ⁻¹¹¹ in relation to a database sequence of human origin were specifically excluded. The final result provided the 2502 sequences listed in the accompanying Sequence Listing. The Sequence Listing is arranged beginning with sequences with no similarity to any sequence in a database searched, and ending with sequences with the greatest similarity. Each identified polynucleotide represents sequence from at least a partial mRNA transcript. Polynucleotides that were determined to be novel were assigned a sequence identification number.

The novel polynucleotides were assigned sequence identification numbers SEQ ID NOS:1-2502. The DNA sequences corresponding to the novel polynucleotides are provided in the Sequence Listing. The majority of the sequences are presented in the Sequence Listing in the 5' to 3' direction. A small number of sequences are listed in the Sequence Listing in the 5' to 3' direction but the sequence as written is actually 3' to 5'. These sequences are readily identified with the designation "AR" in the Sequence Name in Table 1 (inserted before the claims). The sequences correctly listed in the 5' to 3' direction in the Sequence Listing are designated "AF." Table 1 provides: 1) the SEQ ID NO assigned to each sequence for use in the present specification; 2) the filing date of the U.S. priority application in which the sequence was first filed; 3) the SEQ ID NO assigned to the sequence in the priority application; 4) the sequence name used as an internal identifier of

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the sequence; 5) the name assigned to the clone from which the sequence was isolated; and 6) the number of the cluster to which the sequence is assigned (Cluster ID; where the cluster ID is 0, the sequence was not assigned to any cluster

Because the provided polynucleotides represent partial mRNA transcripts, two or more polynucleotides of the invention may represent different regions of the same mRNA transcript and the same gene. Thus, if two or more SEQ ID NOS: are identified as belonging to the same clone, then either sequence can be used to obtain the full-length mRNA or gene. In addition, some sequences are identified with multiple SEQ ID NOS, since these sequences were present in more than one filing. For example, SEQ ID NO:87 and SEQ ID NO:1000 represent the same sequence.

In order to confirm the sequences of SEQ ID NOS:1-2502, inserts of the clones corresponding to these polynucleotides were re-sequenced. These "validation" sequences are provided in SEQ ID NOS:2503-5106. Of these validation sequences, SEQ ID NOS:3040, 3545, 3863, 4511, 4726, and 4749 are not true validation sequences. Instead, SEQ ID NOS:3545, 4511, 4726, and 4749 represent "placeholder" sequences, *i.e.*, sequences that were inserted into the Sequence Listing only to prevent renumbering of the subsequent sequences during generation of the Sequence Listing. Thus, reference to "SEQ ID NOS:1-5252," "SEQ ID NOS:1-5106," or other ranges of SEQ ID NOS that include these placeholder sequences should be read to exclude SEQ ID NOS:3545, 4511, 4726, and 4749.

The validation sequences were often longer than the original polynucleotide sequences they validate, and thus often provide additional sequence information. Validation sequences can be correlated with the original sequences they validate by referring to Table 1. For example, validation sequences of SEQ ID NOS:2503-3039, 3041-3544, 3546-3862 3864-4510, and 4512-4725 share the clone name of the sequence of SEQ ID NOS:1-2502 that they validate.

Example 2: Results of Public Database Search to Identify Function of Gene Products
SEQ ID NOS:1-2502, as well as the validation sequences SEQ ID NOS:2503-3039,
3041-3544, 3546-3862 3864-4510, and 4512-4725 xx:clf were translated in all three
reading frames to determine the best alignment with the individual sequences. These
amino acid sequences and nucleotide sequences are referred, generally, as query sequences,
which are aligned with the individual sequences. Query and individual sequences were

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aligned using the BLAST programs, available over the world wide web at http://ww.ncbi.nlm.nih.gov/BLAST/. Again the sequences were masked to various extents to prevent searching of repetitive sequences or poly-A sequences, using the XBLAST program for masking low complexity as described above in Example 1.

Table 2 (inserted before the claims) shows the results of the alignments. Table 2 refers to each sequence by its SEQ ID NO:, the accession numbers and descriptions of nearest neighbors from the Genbank and Non-Redundant Protein searches, and the p values of the search results.

For each of "SEQ ID NOS:1-5106," the best alignment to a protein or DNA sequence is included in Table 2. The activity of the polypeptide encoded by "SEQ ID NOS:1-5106" is the same or similar to the nearest neighbor reported in Table 2. The accession number of the nearest neighbor is reported, providing a reference to the activities exhibited by the nearest neighbor. The search program and database used for the alignment also are indicated as well as a calculation of the p value.

Full length sequences or fragments of the polynucleotide sequences of the nearest neighbors can be used as probes and primers to identify and isolate the full length sequence of "SEQ ID NOS:1-5106." The nearest neighbors can indicate a tissue or cell type to be used to construct a library for the full-length sequences of "SEQ ID NOS:1-5106."

"SEQ ID NOS:1-5106" and the translations thereof may be human homologs of known genes of other species or novel allelic variants of known human genes. In such cases, these new human sequences are suitable as diagnostics or therapeutics. As diagnostics, the human sequences "SEQ ID NOS:1-5106" exhibit greater specificity in detecting and differentiating human cell lines and types than homologs of other species. The human polypeptides encoded by "SEQ ID NOS1-5106" are likely to be less immunogenic when administered to humans than homologs from other species. Further, on administration to humans, the polypeptides encoded by "SEQ ID NOS:1-5106" can show greater specificity or can be better regulated by other human proteins than are homologs from other species.

Example 3: Members of Protein Families

The validation sequences ("SEQ ID NOS:2503-5106") were used to conduct a profile search as described in the specification above. Several of the polynucleotides of the invention were found to encode polypeptides having characteristics of a polypeptide

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belonging to a known protein families (and thus represent new members of these protein families) and/or comprising a known functional domain (Table 3, inserted prior to claims). Thus the invention encompasses fragments, fusions, and variants of such polynucleotides that retain biological activity associated with the protein family and/or functional domain identified herein.

Start and stop indicate the position within the individual sequences that align with the query sequence having the indicated SEQ ID NO. The direction (Dir) indicates the orientation of the query sequence with respect to the individual sequence, where forward (for) indicates that the alignment is in the same direction (left to right) as the sequence provided in the Sequence Listing and reverse (rev) indicates that the alignment is with a sequence complementary to the sequence provided in the Sequence Listing.

Some polynucleotides exhibited multiple profile hits because, for example, the particular sequence contains overlapping profile regions, and/or the sequence contains two different functional domains. These profile hits are described in more detail below. The acronyms used in Table 3 are provided in parentheses following the full name of the protein family or functional domain to which they refer.

a) Seven Transmembrane Integral Membrane Proteins -- Rhodopsin Family (7tm 1). Several of the validation sequences, and thus their corresponding sequence within SEQ ID NOS:1-2502, correspond to a sequence encoding a polypeptide that is a member of the seven transmembrane receptor rhodopsin family. G-protein coupled receptors of the seven transmembrane rhodopsin family (also called R7G) are an extensive group of hormones, neurotransmitters, and light receptors which transduce extracellular signals by interaction with guanine nucleotide-binding (G) proteins (Strosberg A.D. Eur. J. Biochem. (1991) 196:1, Kerlavage A.R. Curr. Opin. Struct. Biol. (1991) 1:394, Probst, et al., DNA Cell Biol. (1992) 11:1, Savarese, et al., Biochem. J. (1992) 283:1, http://www.gcrdb.uthscsa.edu/, http://swift.embl-heidelberg.de/7tm/. The receptors that are currently known to belong to this family are: 1) 5-hydroxytryptamine (serotonin) 1A to 1F, 2A to 2C, 4, 5A, 5B, 6 and 7 (Branchek T., Curr. Biol. (1993) 3:315); 2) acetylcholine, muscarinic-type, M1 to M5; 3) adenosine A1, A2A, A2B and A3 (Stiles G.L. J. Biol. Chem. (1992) 267:6451; 4) adrenergic alpha-1A to -1C; alpha-2A to -2D; beta-1 to -3 (Friell T. et al., Trends Neurosci. (1988) 11:321); 5) angiotensin II types I and II; 6) bombesin subtypes 3 and 4; 7) bradykinin B1 and B2; 8) c3a and C5a anaphylatoxin; 9) cannabinoid CB1 and CB2; 10) chemokines C-C CC-CKR-1 to CC-CKR-8; 11)

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Chemokines C-X-C CXC-CKR-1 to CXC-CKR-4; 12) Cholecystokinin-A and cholecystokinin-B/gastrin Dopamine D1 to D5 (Stevens C.F., Curr. Biol. (1991) 1:20); 13) Endothelin ET-a and ET-b (Sakurai T. et al., Trends Pharmacol. Sci. (1992) 13:103-107); 14) fMet-Leu-Phe (fMLP) (Nformyl peptide); 15) Follicle stimulating hormone (FSH-R); 16) Galanin; 17) Gastrin-releasing peptide (GRP-R); 18) Gonadotropin-releasing hormone 5 (GNRH-R); 19) Histamine H1 and H2 (gastric receptor I); 20) Lutropinchoriogonadotropic hormone (LSH-R) (Salesse R., et al., Biochimie (1991) 73:109); 21) Melanocortin MC1R to MC5R; 22) Melatonin; 23) Neuromedin B (NMB-R); 24) Neuromedin K (NK-3R); 25) Neuropeptide Y types 1 to 6; 26) Neurotensin (NT-R); 27) Octopamine (tyramine), from insects; 28) Odorants (Lancet D., et al., Curr. Biol. 10 (1993)3:668: 29) Opioids delta-, kappa- and mu-types (Uhl G.R., et al., Trends Neurosci. (1994) 17:89; 30) Oxytocin (OT-R); 31) Platelet activating factor (PAF-R); 32) Prostacyclin; 33) Prostaglandin D2; 34) Prostaglandin E2, EP1 to EP4 subtypes; 35) Prostaglandin F2; 36) Purinoreceptors (ATP) (Barnard E.A., et al., Trends Pharmacol. Sci. (1994)15:67; 37); Somatostatin types 1 to 5; 38) Substance-K (NK-2R); Substance-P (NK-15 1R); 39) Thrombin; 40) Thromboxane A2; 41) Thyrotropin (TSH-R) (Salesse R., et al., Biochimie (1991) 73:109); 42) Thyrotropin releasing factor (TRH-R); 42) Vasopressin V1a, V1b and V2; 43) Visual pigments (opsins and rhodopsin) (Applebury M.L., et al., Vision Res. (1986) 26:1881; 44) Proto-oncogene mas; 45) A number of orphan receptors (whose ligand is not known) from mammals and birds; 46) Caenorhabditis elegans putative 20 receptors C06G4.5, C38C10.1, C43C3.2; 47) T27D1.3 and ZC84.4; 48) Three putative

The structure of these receptors is thought to be identical. They have seven hydrophobic regions, each of which most probably spans the membrane. The N-terminus is located on the extracellular side of the membrane and is often glycosylated, while the C-terminus is cytoplasmic and generally phosphorylated. Three extracellular loops alternate with three intracellular loops to link the seven transmembrane regions. Most, but not all of these receptors, lack a signal peptide. The most conserved parts of these proteins are the transmembrane regions and the first two cytoplasmic loops. A conserved acidic-Argaromatic triplet is present in the N-terminal extremity of the second cytoplasmic loop (Attwood T.K., Eliopoulos E.E., Findlay J.B.C. *Gene* (1991) 98:153-159) and could be implicated in the interaction with G proteins.

receptors encoded in the genome of cytomegalovirus: US27, US28, and UL33; and 49)

ECRF3, a putative receptor encoded in the genome of herpesvirus saimiri.

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A consensus pattern that contains the conserved triplet and that also spans the major part of the third transmembrane helix is used to detect this widespread family of proteins: [GSTALIVMFYWC]-[GSTANCPDE]-{EDPKRH}-x(2)-[LIVMNQGA]-x(2)- [LIVMFT]-[GSTANC]-[LIVMFYWSTAC]-[DENH]-R-[FYWCSH]-x(2)- [LIVM].

b) Seven Transmembrane Integral Membrane Proteins -- Secretin Family (7tm 2). Several of the validation sequences, and thus their corresponding sequence within SEQ ID NOS:1-2502, correspond to a sequence encoding a polypeptide that is a member of the seven transmembrane receptor secretin family. A number of peptide hormones bind to Gprotein coupled receptors that, while structurally similar to the majority of G-protein coupled receptors (R7G) (see profile for 7 transmembrane receptors (rhodopsin family), do not show any similarity at the level of their sequence, thus new family whose current known members (Jueppner et al. Science (1991) 254:1024; Hamann et al. Genomics (1996) 32:144).are: 1) calcitonin receptor, 2) calcitonin gene-related peptide receptor; 3) corticotropin releasing factor receptor types 1 and 2; 4) gastric inhibitory polypeptide receptor; 5) glucagon receptor; 6) glucagon-like peptide 1 receptor; 7) growth hormonereleasing hormone receptor; 7) parathyroid hormone / parathyroid hormone-related peptide types 1 and 2; 8) pituitary adenylate cyclase activating polypeptide receptor; 9) secretin receptor; 10) vasoactive intestinal peptide receptor types 1 and 2; 10) insects diuretic hormone receptor; 11) Caenorhabditis elegans putative receptor C13B9.4; 12) Caenorhabditis elegans putative receptor ZK643.3; 13) human leucocyte CD97 (which contains 3 EGF-like domains in its N-terminal section); 14) human cell surface

contains 3 EGF-like domains in its N-terminal section); 14) human cell surface glycoprotein EMR1 (which contains 6 EGF-like domains in it N-terminal section); and 15) mouse cell surface glycoprotein F4/80 (which contains 7 EGF-like domains in its N-terminal section). All of 1) through 10) are coupled to G-proteins which activate both adenylyl cyclase and the phosphatidylinositol-calcium pathway.

Like classical R7G the secretin family of 7 transmembrane proteins contain seven transmembrane regions. Their N-terminus is located on the extracellular side of the membrane and potentially glycosylated, while their C-terminus is cytoplasmic. But apart from these topological similarities they do not share any region of sequence similarity and are therefore probably not evolutionary related.

Every receptor in the 7 transmember secretin family is encoded on multiple exons, and several of these functionally distinct products. The N-terminal extracellular domain of these receptors contains five conserved cysteines residues that may be involved in disulfide

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bonds, with a consensus pattern in the region that spans the first three cysteines. One of the most highly conserved regions spans the C-terminal part of the last transmembrane region and the beginning of the adjacent intracellular region. This second region is used as a second signature pattern. The two consensus patterns are:

- 1) C-x(3)-[FYWLIV]-D-x(3,4)-C-[FW]-x(2)-[STAGV]-x(8,9)-C-[PF]
 2) Q-G-[LMFCA]-[LIVMFT]-[LIV]-x-[LIVFST]-[LIF]-[VFYH]-C- [LFY]-x-N-x(2)-V
 - c) Ank Repeats (ANK). SEQ IS NO:2656, and thus its corresponding sequence within SEQ ID NOS:1-2502, represents a polynucleotide encoding an Ank repeat-containing protein. The ankyrin motif is a 33 amino acid sequence named after the protein ankyrin which has 24 tandem 33-amino-acid motifs. Ank repeats were originally identified in the cell-cycle-control protein cdc10 (Breeden et al., Nature (1987) 329:651). Proteins containing ankyrin repeats include ankyrin, myotropin, I-kappaB proteins, cell cycle protein cdc10, the Notch receptor (Matsuno et al., Development (1997) 124(21):4265); G9a (or BAT8) of the class III region of the major histocompatibility complex (Biochem J. 290:811-818, 1993), FABP, GABP, 53BP2, Lin12, glp-1, SW14, and SW16. The functions of the ankyrin repeats are compatible with a role in protein-protein interactions (Bork, Proteins (1993) 17(4):363; Lambert and Bennet, Eur. J. Biochem. (1993) 211:1; Kerr et al., Current Op. Cell Biol. (1992) 4:496; Bennet et al., J. Biol. Chem. (1980) 255:6424).

The 90 kD N-terminal domain of ankyrin contains a series of 24 33-amino-acid ank repeats. (Lux et al., Nature (1990) 344:36-42, Lambert et al., PNAS USA (1990) 87:1730.) The 24 ank repeats form four folded subdomains of 6 repeats each. These four repeat subdomains mediate interactions with at least 7 different families of membrane proteins. Ankyrin contains two separate binding sites for anion exchanger dimers. One site utilizes repeat subdomain two (repeats 7-12) and the other requires both repeat subdomains 3 and 4 (repeats 13-24). Since the anion exchangers exist in dimers, ankyrin binds 4 anion exchangers at the same time (Michaely and Bennett, J. Biol. Chem. (1995) 270(37):22050). The repeat motifs are involved in ankyrin interaction with tubulin, spectrin, and other membrane proteins. (Lux et al., Nature (1990) 344:36.)

The Rel/NF-kappaB/Dorsal family of transcription factors have activity that is controlled by sequestration in the cytoplasm in association with inhibitory proteins referred to as I-kappaB. (Gilmore, Cell (1990) 62:841; Nolan and Baltimore, Curr Opin Genet Dev. (1992) 2:211; Baeuerle, Biochim Biophys Acta (1991) 1072:63; Schmitz et al., Trends Cell

Biol. (1991) 1:130.) I-kappaB proteins contain 5 to 8 copies of 33 amino acid ankyrin repeats and certain NF-kappaB/rel proteins are also regulated by cis-acting ankyrin repeat containing domains including p105NF-kappaB which contains a series of ankyrin repeats (Diehl and Hannink, J. Virol. (1993) 67(12):7161). The I-kappaBs and Cactus (also containing ankyrin repeats) inhibit activators through differential interactions with the Relhomology domain. The gene family includes proto-oncogenes, thus broadly implicating I-kappaB in the control of both normal gene expression and the aberrant gene expression that makes cells cancerous. (Nolan and Baltimore, Curr Opin Genet Dev. (1992) 2(2):211-220). In the case of rel/NF-kappaB and pp40/I-kappaB(, both the ankyrin repeats and the carboxy-terminal domain are required for inhibiting DNA-binding activity and direct association of pp40/I-kappaB(with rel/NF-kappaB protein. The ankyrin repeats and the carboxy-terminal of pp40/I-kappaB(form a structure that associates with the rel homology domain to inhibit DNA binding activity (Inoue et al., PNAS USA (1992) 89:4333).

The 4 ankyrin repeats in the amino terminus of the transcription factor subunit GABP—are required for its interaction with the GABP—subunit to form a functional high affinity DNA-binding protein. These repeats can be crosslinked to DNA when GABP is bound to its target sequence. (Thompson et al., Science (1991) 253:762; LaMarco et al., Science (1991) 253:789). Myotrophin, a 12.5 kDa protein having a key role in the initiation of cardiac hypertrophy, comprises ankyrin repeats. The ankyrin repeats are characteristic of a hairpin-like protruding tip followed by a helix-turn-helix motif. The V-shaped helix-turn-helix of the repeats stack sequentially in bundles and are stabilized by compact hydrophobic cores, whereas the protruding tips are less ordered.

d) Eukaryotic Aspartyl Proteases (asp). Several of the validation sequences, and thus their corresponding sequence within SEQ ID NOS:1-2502, correspond to a sequence encoding a novel eukaryotic aspartyl protease. Aspartyl proteases, known as acid proteases, (EC 3.4.23.-) are a widely distributed family of proteolytic enzymes (Foltmann B., Essays Biochem. (1981) 17:52; Davies D.R., Annu. Rev. Biophys. Chem. (1990) 19:189; Rao J.K.M., et al., Biochemistry (1991) 30:4663) known to exist in vertebrates, fungi, plants, retroviruses and some plant viruses. Aspartate proteases of eukaryotes are monomeric enzymes which consist of two domains. Each domain contains an active site centered on a catalytic aspartyl residue. The two domains most probably evolved from the duplication of an ancestral gene encoding a primordial domain. Currently known eukaryotic aspartyl proteases include: 1) Vertebrate gastric pepsins A and C (also known as

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gastricsin); 2) Vertebrate chymosin (rennin), involved in digestion and used for making cheese; 3) Vertebrate lysosomal cathepsins D (EC 3.4.23.5) and E (EC 3.4.23.34); 4) Mammalian renin (EC 3.4.23.15) whose function is to generate angiotensin I from angiotensinogen in the plasma; 5) Fungal proteases such as aspergillopepsin A (EC 3.4.23.18), candidapepsin (EC 3.4.23.24), mucoropepsin (EC 3.4.23.23) (mucor rennin), endothiapepsin (EC 3.4.23.22), polyporopepsin (EC 3.4.23.29), and rhizopuspepsin (EC 3.4.23.21); and 6) Yeast saccharopepsin (EC 3.4.23.25) (proteinase A) (gene PEP4). PEP4 is implicated in posttranslational regulation of vacuolar hydrolases; 7) Yeast barrierpepsin (EC 3.4.23.35) (gene BAR1); a protease that cleaves alpha-factor and thus acts as an antagonist of the mating pheromone; and 8) Fission yeast sxal which is involved in 10 degrading or processing the mating pheromones.

Most retroviruses and some plant viruses, such as badnaviruses, encode for an aspartyl protease which is an homodimer of a chain of about 95 to 125 amino acids. In most retroviruses, the protease is encoded as a segment of a polyprotein which is cleaved during the maturation process of the virus. It is generally part of the pol polyprotein and, more rarely, of the gag polyprotein. Because the sequence around the two aspartates of eukaryotic aspartyl proteases and around the single active site of the viral proteases is conserved, a single signature pattern can be used to identify members of both groups of proteases. The consensus pattern is: [LIVMFGAC]-[LIVMTADN]-[LIVFSA]-D-[ST]-G-[STAV]-[STAPDENQ]- x-[LIVMFSTNC]-x-[LIVMFGTA], where D is the active site residue.

e) ATPases Associated with Various Cellular Activities (ATPases). Several of the validation sequences, and thus their corresponding sequence within SEQ ID NOS:1-2502, correspond to a sequence that encodes a novel member of the "ATPases Associated with diverse cellular Activities" (AAA) protein family. The AAA protein family is composed of a large number of ATPases that share a conserved region of about 220 amino acids that contains an ATP-binding site (Froehlich et al., J. Cell Biol. (1991) 114:443; Erdmann et al. Cell (1991) 64:499; Peters et al., EMBO J. (1990) 9:1757; Kunau et al., Biochimie (1993) 75:209-224; Confalonieri et al., BioEssays (1995) 17:639; http://yeamob.pci.chemie.unituebingen.de/AAA/Description.html). The proteins that belong to this family either contain one or two AAA domains.

Proteins containing two AAA domains include: 1) Mammalian and drosophila NSF (N-ethylmaleimide-sensitive fusion protein) and the fungal homolog, SEC18, which are

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involved in intracellular transport between the endoplasmic reticulum and Golgi, as well as between different Golgi cisternae; 2) Mammalian transitional endoplasmic reticulum ATPase (previously known as p97 or VCP), which is involved in the transfer of membranes from the endoplasmic reticulum to the golgi apparatus. This ATPase forms a ring-shaped homooligomer composed of six subunits. The yeast homolog, CDC48, plays a role in spindle pole proliferation; 3) Yeast protein PAS1 essential for peroxisome assembly and the related protein PAS1 from Pichia pastoris; 4) Yeast protein AFG2; 5) Sulfolobus acidocaldarius protein SAV and Halobacterium salinarium cdcH, which may be part of a transduction pathway connecting light to cell division.

Proteins containing a single AAA domain include: 1) Escherichia coli and other bacteria ftsH (or hflB) protein. FtsH is an ATP-dependent zinc metallopeptidase that degrades the heat-shock sigma-32 factor, and is an integral membrane protein with a large cytoplasmic C-terminal domain that contain both the AAA and the protease domains; 2) Yeast protein YME1, a protein important for maintaining the integrity of the mitochondrial compartment. YME1 is also a zinc-dependent protease; 3) Yeast protein AFG3 (or YTA10). This protein also contains an AAA domain followed by a zinc-dependent protease domain; 4) Subunits from regulatory complex of the 26S proteasome (Hilt et al., Trends Biochem. Sci. (1996) 21:96), which is involved in the ATP-dependent degradation of ubiquitinated proteins, which subunits include: a) Mammalian 4 and homologs in other higher eukaryotes, in yeast (gene YTA5) and fission yeast (gene mts2); b) Mammalian 6 (TBP7) and homologs in other higher eukaryotes and in yeast (gene YTA2); c) Mammalian subunit 7 (MSS1) and homologs in other higher eukaryotes and in yeast (gene CIM5 or YTA3); d) Mammalian subunit 8 (P45) and homologs in other higher eukaryotes and in yeast (SUG1 or CIM3 or TBY1) and fission yeast (gene let1); e) Other probable subunits include human TBP1, which influences HIV gene expression by interacting with the virus tat transactivator protein, and yeast YTA1 and YTA6; 5) Yeast protein BCS1, a mitochondrial protein essential for the expression of the Rieske iron-sulfur protein; 6) Yeast protein MSP1, a protein involved in intramitochondrial sorting of proteins; 7) Yeast protein PAS8, and the corresponding proteins PAS5 from Pichia pastoris and PAY4 from Yarrowia lipolytica; 8) Mouse protein SKD1 and its fission yeast homolog (SpAC2G11.06); 9) Caenorhabditis elegans meiotic spindle formation protein mei-1; 10) Yeast protein SAP1' 11) Yeast protein YTA7; and 12) Mycobacterium leprae hypothetical protein A2126A.

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In general, the AAA domains in these proteins act as ATP-dependent protein clamps(Confalonieri *et al.* (1995) *BioEssays 17*:639). In addition to the ATP-binding 'A' and 'B' motifs, which are located in the N-terminal half of this domain, there is a highly conserved region located in the central part of the domain which was used in the development of the signature pattern. The consensus pattern is: [LIVMT]-x-[LIVMT]-[LIVMT]-x-[GATMC]-[ST]-[NS]-x(4)-[LIVM]- D-x-A-[LIFA]-x-R.

f) <u>Bcl-2 family (Bcl-2)</u>. SEQ ID NO:3404, and thus the corresponding sequence it validates, represents a polynucleotide encoding an apoptosis regulator protein of the Bcl-2 family. Active cell suicide (apoptosis) is induced by events such as growth factor withdrawal and toxins. It is controlled by regulators, which have either an inhibitory effect on programmed cell death (anti-apoptotic) or block the protective effect of inhibitors (proapoptotic) (Vaux, 1993, Curr. Biol. 3:877-878, and White, 1996, Genes Dev. 10:2859-2869). Many viruses have found a way of countering defensive apoptosis by encoding their own anti-apoptosis genes, preventing their target cells from dying prematurely.

All proteins belonging to the Bcl-2 family (Reed et al., 1996, Adv. Exp. Med. Biol. 406:99-112) contain either a BH1, BH2, BH3, or BH4 domain. All anti-apoptotic proteins contain BH1 and BH2 domains; some of them contain an additional N-terminal BH4 domain (Bcl-2, Bcl-x(L), Bcl-w), which is never seen in pro-apoptotic proteins, except for Bcl-x(S). On the other hand, all pro-apoptotic proteins contain a BH3 domain (except for Bad) necessary for dimerization with other proteins of Bcl-2 family and crucial for their killing activity; some of them also contain BH1 and BH2 domains (Bax, Bak). The BH3 domain is also present in some anti-apoptotic protein, such as Bcl-2 or Bcl-x(L). Proteins that are known to contain these domains are listed below.

- 1. Vertebrate protein Bcl-2. Bcl-2 blocks apoptosis; it prolongs the survival of hematopoietic cells in the absence of required growth factors and also in the presence of various stimuli inducing cellular death. Two isoforms of bcl-2 (alpha and beta) are generated by alternative splicing. Bcl-2 is expressed in a wide range of tissues at various times during development. It forms heterodimers with the Bax proteins.
- 2. Vertebrate protein Bcl-x. Two isoforms of Bcl-x (Bcl-x(L) and Bcl-x(S)) are generated by alternative splicing. While the longer product (Bcl-x(L)) can protect a growth-factor-dependent cell line from apoptosis, the shorter form blocks the protective effect of Bcl-2 and Bcl-x(L) and acts as an anti-anti-apoptosis protein.
- 3. Mammalian protein Bax. Bax blocks the anti-apoptosis ability of Bcl-2 with which

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it forms heterodimers. There is no evidence that Bax has any activity in the absence of Bcl-2. Three isoforms of bax (alpha, beta and gamma) are generated by alternative splicing.

- 4. Mammalian protein Bak, which promotes cell death and counteracts the protection from apoptosis provided by Bcl-2.
- 5. Mammalian protein Bcl-w, which promotes cell survival.
- 6. Mammalian protein bad, which promotes cell death, and counteracts the protection from apoptosis provided by Bcl-x(L), but not that of Bcl-2.
- 7. Human protein Bik, which promotes cell death, but cannot counteract the protection from apoptosis provided by Bcl-2.
 - 8. Mouse protein Bid, which induces caspases and apoptosis, and counteracts the protection from apoptosis provided by Bcl-2.
 - 9. Human induced myeloid leukemia cell differentiation protein MCL1. MCL1 is probably involved in programming of differentiation and concomitant maintenance of viability but not proliferation. Its expression increases early during phorbol ester induced differentiation in myeloid leukemia cell line ML-1.
 - 10. Mouse hemopoietic-specific early response protein A1.
 - 11. Mammalian activator of apoptosis Harakiri (Inohara et al., 1997, EMBO J. 16:1686-1694) (also known as neuronal death protein Dp5). This is a small protein of 92 residues that activates apoptosis. It contains a BH3 domain, but no BH1, BH2 or BH4 domains.

The following consensus patterns have been developed for the four BH domains:

- 1) [LVME]-[FT]-x-[GSD]-[GL]-x(1,2)-[NS]-[YW]-G-R-[LIV]- [LIVC]-[GAT]-[LIVMF](2)-x-F-[GSAE]-[GSARY]
- 25 2) W-[LIM]-x(3)-[GR]-G-[WQ]-[DENSAV]-x-[FLGA]-[LIVFTC]
 - 3) [LIVAT]-x(3)-L-[KARQ]-x-[IVAL]-G-D-[DESG]-[LIMFV]-[DENSHQ]-[LVSHRQ]-[NSR]
 - 4) [DS]-[NT]-R-[AE]-[LI]-V-x-[KD]-[FY]-[LIV]-[GHS]-Y-K-L- [SR]-Q-[RK]-G-[HY]-x-[CW].
- g) <u>Bromodomain (bromodomain)</u>. SEQ ID NOS:4036 and 4489, and thus the corresponding sequences they validate, represent polynucleotides encoding a polypeptide having a bromodomain region (Haynes et al., 1992, Nucleic Acids Res. 20:2693-2603, Tamkun et al., 1992, Cell 68:561-572, and Tamkun, 1995, Curr. Opin. Genet. Dev. 5:473-

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477), which is a conserved region of about 70 amino acids found in the following proteins: 1) Higher eukaryotes transcription initiation factor TFIID 250 Kd subunit (TBP-associated factor p250) (gene CCG1); P250 is associated with the TFIID TATA-box binding protein and seems essential for progression of the G1 phase of the cell cycle. 2) Human RING3, a protein of unknown function encoded in the MHC class II locus; 3) Mammalian CREB-binding protein (CBP), which mediates cAMP-gene regulation by binding specifically to phosphorylated CREB protein; 4) Mammalian homologs of brahma, including three brahma-like human: SNF2a(hBRM), SNF2b, and BRG1; 5) Human BS69, a protein that binds to adenovirus E1A and inhibits E1A transactivation; 6) Human peregrin (or Br140).

The bromodomain is thought to be involved in protein-protein interactions and may be important for the assembly or activity of multicomponent complexes involved in transcriptional activation. The consensus pattern, which spans a major part of the bromodomain, is: [STANVF]-x(2)-F-x(4)-[DNS]-x(5,7)-[DENQTF]-Y-[HFY]-x(2)-[LIVMFY]-x(3)-[LIVM]-x(4)-[LIVM]-x(6,8)-Y-x(12,13)-[LIVM]-x(2)-N-[SACF]-x(2)-[FY].

- h) Basic Region Plus Leucine Zipper Transcription Factors (BZIP). SEQ ID NO:3408, 2951, and 4850, and thus the corresponding sequences these sequences validate, represent polynucleotides encoding a novel member of the family of basic region plus leucine zipper transcription factors. The bZIP superfamily (Hurst, Protein Prof. (1995) 2:105; and Ellenberger, Curr. Opin. Struct. Biol. (1994) 4:12) of eukaryotic DNA-binding transcription factors encompasses proteins that contain a basic region mediating sequencespecific DNA-binding followed by a leucine zipper required for dimerization. Members of the family include transcription factor AP-1, which binds selectively to enhancer elements in the cis control regions of SV40 and metallothionein IIA. AP-1, also known as c-jun, is the cellular homolog of the avian sarcoma virus 17 (ASV17) oncogene v-jun. Other members of this protein family include jun-B and jun-D, probable transcription factors that are highly similar to jun/AP-1; the fos protein, a proto-oncogene that forms a non-covalent dimer with c-jun; the fos-related proteins fra-1, and fos B; and mammalian cAMP response element (CRE) binding proteins CREB, CREM, ATF-1, ATF-3, ATF-4, ATF-5, ATF-6 and LRF-1. The consensus pattern for this protein family is: [KR]-x(1,3)-[RKSAQ]-N-x(2)-[SAQ](2)-x-[RKTAENQ]-x-R-x-[RK].
- i) Cyclins (cyclin). SEQ ID NOS:3618, 3895, and 4536, and thus the corresponding sequences these sequences validate, represent polynucleotides encoding

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cyclins, and SEQ ID NO:55 and 56, respectively, show the corresponding full-length polynucleotides. SEQ ID NO:57 and 58 show, respectively, the translations of SEQ ID NO:55 and 56. Cyclins (Nurse, 1990, Nature 344:503-508; Norbury et al., 1991, Curr. Biol. 1:23-24; and Lew et al., 1992, Trends Cell Biol. 2:77-81) are eukaryotic proteins that play an active role in controlling nuclear cell division cycles. There are two main groups of cyclins. G2/M cyclins are essential for the control of the cell cycle at the G2/M (mitosis) transition. G2/M cyclins accumulate steadily during G2 and are abruptly destroyed as cells exit from mitosis (at the end of the M-phase). G1/S cyclins are essential for the control of the cell cycle at the G1/S (start) transition.

The best conserved region is in the central part of the cyclins' sequences, known as the "cyclin-box," from which a 32 residue consensus pattern was derived: R-x(2)-[LIVMSA]-x(2)-[FYWS]-[LIVM]-x(8)-[LIVMFC]-x(4)-[LIVMFYA]-x(2)-[STAGC]-[LIVMFYQ]-x-[LIVMFYC]-[LIVMFY]-D-[RKH]-[LIVMFYW].

j) Eukaryotic thiol (cysteine) proteases active sites (Cys-protease). SEQ ID NOS:3344, 3684, 3688, and 4801, and thus also the sequences they validate, represent polynucleotides encoding proteins having a eukaryotic thiol (cysteine) protease active site. Eukaryotic thiol proteases (Dufour E., Biochimie (1988) 70:1335); are a family of proteolytic enzymes which contain an active site cysteine. Catalysis proceeds through a thioester intermediate and is facilitated by a nearby histidine side chain; an asparagine completes the essential catalytic triad. The proteases that belong to this family are: 1) vertebrate lysosomal cathepsins B (Kirschke H., et al., *Protein Prof.* (1995) 2:1587-1643); 2) vertebrate lysosomal dipeptidyl peptidase I (also known as cathepsin C) (Kirschke H., et al., supra); 3) vertebrate calpains (Calpains are intracellular calciumactivated thiol protease that contain both an N-terminal catalytic domain and a C-terminal calcium-binding domain); 4) mammalian cathepsin K, which seems involved in osteoclastic bone resorption (Shi G.-P., et al., FEBS Lett. (1995) 357:129); 5) human cathepsin O ([4] Velasco G., Ferrando A.A., Puente X.S., Sanchez L.M., Lopez-Otin C. J. Biol. Chem. (1994) 269:27136); 6) bleomycin hydrolase (which catalyzes the inactivation of the antitumor drug BLM (a glycopeptide)); 7) Plant enzymes such as: barley aleurain, EP-B1/B4; kidney bean EP-C1, rice bean SH-EP; kiwi fruit actinidin; papaya latex papin, chymopapain, caricain, and proteinase IV; pea turgor-responsive protein 15A; pineapple stem bromelain; rape COT44; rice oryzain alpha, beta, and gamma; tomato lowtemperature induced, Arabidopsis thaliana A494, RD19A and RD21A; 8) - House-dust

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mites allergens DerP1 and EurM1; 9) cathepsin B-like proteinases from the worms

Caenorhabditis elegans (genes gcp-1, cpr-3, cpr-4, cpr-5 and cpr-6), Schistosoma mansoni
(antigen SM31) and Japonica (antigen SJ31), Haemonchus contortus (genes AC-1 and
AC-2), and Ostertagia ostertagi (CP-1 and CP-3); 10) slime mold cysteine proteinases CP1
and CP2; 11) cruzipain from Trypanosoma cruzi and brucei; 12) throphozoite cysteine
proteinase (TCP) from various Plasmodium species; 13) proteases from Leishmania
mexicana, Theileria annulata and Theileria parva; 14) Baculoviruses cathepsin-like enzyme
(v-cath); 15) Drosophila small optic lobes protein (gene sol), a neuronal protein that
contains a calpain-like domain; 16) yeast thiol protease BLH1/YCP1/LAP3;

17) Caenorhabditis elegans hypothetical protein C06G4.2, a calpain-like protein.

In addition, two bacterial peptidases are also part of this family: 1) aminopeptidase C from Lactococcus lactis (gene pepC) (Chapot-Chartier M.P., et al., *Appl. Environ. Microbiol.* (1993) 59:330); and 2) thiol protease tpr from Porphyromonas gingivalis. Three other proteins are structurally related to this family, but may have lost their proteolytic activity. These include: 1) soybean oil body protein P34 (which has its active site cysteine replaced by a glycine); 2) rat testin (which is a sertoli cell secretory protein highly similar to cathepsin L but with the active site cysteine is replaced by a serine); and 3) Plasmodium falciparum serine-repeat protein (SERA) (which is the major blood stage antigen and possesses a C-terminal thiol-protease-like domain (Higgins D.G., et al., *Nature* (1989) 340:604), with the active site cysteine is replaced by a serine).

The sequences around the three active site residues are well conserved and can be used as signature patterns:

Consensus pattern #1: Q-x(3)-[GE]-x-C-[YW]-x(2)-[STAGC]-[STAGCV] (where C is the active site residue)

Consensus pattern #2: [LIVMGSTAN]-x-H-[GSACE]-[LIVM]-x-[LIVMAT](2)-G-x-[GSADNH] (where H is the active site residue);

Consensus patern #3: [FYCH]-[WI]-[LIVT]-x-[KRQAG]-N-[ST]-W-x(3)-[FYW]-G-x(2)-G- [LFYW]-[LIVMFYG]-x-[LIVMF] (where N is the active site residue).

k) Phorbol Esters/Diacylglycerol Binding (DAG_PE_bind). SEQ ID NO:4659, and thus the sequence it validates, represents a polynucleotide encoding a protein belonging to the family including phorbol esters/diacylglycerol binding proteins. Diacylglycerol (DAG) is an important second messenger. Phorbol esters (PE) are analogues of DAG and potent tumor promoters that cause a variety of physiological changes when administered to both

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cells and tissues. DAG activates a family of serine/threonine protein kinases, collectively known as protein kinase C (PKC) (Azzi et al., Eur. J. Biochem. (1992) 208:547). Phorbol esters can directly stimulate PKC. The N-terminal region of PKC, known as C1, has been shown (Ono et al., Proc. Natl. Acad. Sci. USA (1989) 86:4868) to bind PE and DAG in a phospholipid and zinc-dependent fashion. The C1 region contains one or two copies (depending on the isozyme of PKC) of a cysteine-rich domain about 50 amino-acid residues long and essential for DAG/PE-binding. Such a domain has also been found in, for example, the following proteins.

- (1) Diacylglycerol kinase (EC 2.7.1.107) (DGK) (Sakane *et al.*, *Nature* (1990) 344:345), the enzyme that converts DAG into phosphatidate. It contains two copies of the DAG/PE-binding domain in its N-terminal section. At least five different forms of DGK are known in mammals; and
- (2) N-chimaerin, a brain specific protein which shows sequence similarities with the BCR protein at its C-terminal part and contains a single copy of the DAG/PE-binding domain at its N-terminal part. It has been shown (Ahmed *et al.*, *Biochem. J.* (1990) 272:767, and Ahmed *et al.*, *Biochem. J.* (1991) 280:233) to be able to bind phorbol esters.

The DAG/PE-binding domain binds two zinc ions; the ligands of these metal ions are probably the six cysteines and two histidines that are conserved in this domain. The signature pattern completely spans the DAG/PE domain. The consensus pattern is: H-x-[LIVMFYW]-x(8,11)-C-x(2)-C-x(3)-[LIVMFC]-x(5,10)-C-x(2)-C-x(4)-[HD]-x(2)-C-x(5,9)-C. All the C and H are probably involved in binding zinc.

- 1) <u>DEAD and DEAH box families ATP-dependent helicases signatures</u>
 (<u>Dead_box_helic</u>). SEQ ID NOS:4821 and 5083, and thus the sequences they validate, represent polynucleotides encoding a novel member of the DEAD box family. A number of eukaryotic and prokaryotic proteins have been characterized (Schmid S.R., et al., *Mol. Microbiol.* (1992) 6:283; Linder P., et al., *Nature* (1989) 337:121; Wassarman D.A., et al., *Nature* (1991) 349:463) on the basis of their structural similarity. All are involved in ATP-dependent, nucleic-acid unwinding. Proteins currently known to belong to this family are:
- 1) Initiation factor eIF-4A. Found in eukaryotes, this protein is a subunit of a high molecular weight complex involved in 5'cap recognition and the binding of mRNA to ribosomes. It is an ATP-dependent RNA-helicase.
 - 2) PRP5 and PRP28. These yeast proteins are involved in various ATP-requiring steps of the pre-mRNA splicing process.

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- 3) Pl10, a mouse protein expressed specifically during spermatogenesis.
- 4) An3, a Xenopus putative RNA helicase, closely related to Pl10.
- 5) SPP81/DED1 and DBP1, two yeast proteins involved in pre-mRNA splicing and related to Pl10.
- 6) Caenorhabditis elegans helicase glh-1.
 - 7) MSS116, a yeast protein required for mitochondrial splicing.
 - 8) SPB4, a yeast protein involved in the maturation of 25S ribosomal RNA.
 - 9) p68, a human nuclear antigen. p68 has ATPase and DNA-helicase activities in vitro. It is involved in cell growth and division.
- 10 10) Rm62 (p62), a Drosophila putative RNA helicase related to p68.
 - 11) DBP2, a yeast protein related to p68.
 - 12) DHH1, a yeast protein.
 - 13) DRS1, a yeast protein involved in ribosome assembly.
 - 14) MAK5, a yeast protein involved in maintenance of dsRNA killer plasmid.
- 15) ROK1, a yeast protein.
 - 16) ste13, a fission yeast protein.
 - 17) Vasa, a Drosophila protein important for oocyte formation and specification of embryonic posterior structures.
 - 18) Me31B, a Drosophila maternally expressed protein of unknown function.
 - 19) dbpA, an Escherichia coli putative RNA helicase.
 - 20) deaD, an Escherichia coli putative RNA helicase which can suppress a mutation in the rpsB gene for ribosomal protein S2.
 - 21) rhlB, an Escherichia coli putative RNA helicase.
 - 22) rhlE, an Escherichia coli putative RNA helicase.
- 25 23) rmB, an Escherichia coli protein that shows RNA-dependent ATPase activity, which interacts with 23S ribosomal RNA.
 - 24) Caenorhabditis elegans hypothetical proteins T26G10.1, ZK512.2 and ZK686.2.
 - 25) Yeast hypothetical protein YHR065c.
- 30 26) Yeast hypothetical protein YHR169w.
 - 27) Fission yeast hypothetical protein SpAC31A2.07c.
 - 28) Bacillus subtilis hypothetical protein yxiN.

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All of the above proteins share a number of conserved sequence motifs. Some of them are specific to this family while others are shared by other ATP-binding proteins or by proteins belonging to the helicases 'superfamily' (Hodgman T.C., *Nature* (1988) 333:22 and *Nature* (1988) 333:578 (Errata);

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- http://www.expasy.ch/www/linder/HELICASES_TEXT.html). One of these motifs, called the 'D-E-A-D-box', represents a special version of the B motif of ATP-binding proteins.
 Some other proteins belong to a subfamily which have His instead of the second Asp and are thus said to be 'D-E-A-H-box' proteins (Wassarman D.A., et al., *Nature* (1991) 349:463; Harosh I., et al., *Nucleic Acids Res.* (1991) 19:6331; Koonin E.V., et al., *J. Gen.* Virol. (1992) 73:989; http://www.expasy.ch/www/linder/HELICASES_TEXT.html).
 Proteins currently known to belong to this DEAH subfamily are:
 - 1) PRP2, PRP16, PRP22 and PRP43. These yeast proteins are all involved in various ATP-requiring steps of the pre-mRNA splicing process. 2) Fission yeast prh1, which my be involved in pre-mRNA splicing. 3) Male-less (mle), a Drosophila protein required in males, for dosage compensation of X chromosome linked genes. 4) RAD3 from yeast. RAD3 is a DNA helicase involved in excision repair of DNA damaged by UV light, bulky adducts or cross-linking agents. Fission yeast rad15 (rhp3) and mammalian DNA excision repair protein XPD (ERCC-2) are the homologs of RAD3. 5) Yeast CHL1 (or CTF1), which is important for chromosome transmission and normal cell cycle progression in G(2)/M. 6) Yeast TPS1. 7) Yeast hypothetical protein YKL078w. 8) Caenorhabditis elegans hypothetical proteins C06E1.10 and K03H1.2. 9) Poxviruses' early transcription factor 70 Kd subunit which acts with RNA polymerase to initiate transcription from early gene promoters. 10) 18, a putative vaccinia virus helicase. 11) hrpA, an Escherichia coli putative RNA helicase.

The following signature patterns are used to identify member for both subfamilies: Consensus pattern: [LIVMF](2)-D-E-A-D-[RKEN]-x-[LIVMFYGSTN]
Consensus pattern: [GSAH]-x-[LIVMF](3)-D-E-[ALIV]-H-[NECR].

m) <u>EF Hand (EFhand)</u>. Several of the validation sequences, and thus the sequences they validate, correspond to polynucleotides encoding a novel protein in the family of EFhand proteins. Many calcium-binding proteins belong to the same evolutionary family and share a type of calcium-binding domain known as the EF-hand (Kawasaki *et al.*, *Protein*. *Prof.* (1995) 2:305-490). This type of domain consists of a twelve residue loop flanked on both sides by a twelve residue alpha-helical domain. In an EF-hand loop the calcium ion is

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[LIVMFYW].

coordinated in a pentagonal bipyramidal configuration. The six residues involved in the binding are in positions 1, 3, 5, 7, 9 and 12; these residues are denoted by X, Y, Z, -Y, -X and -Z. The invariant Glu or Asp at position 12 provides two oxygens for liganding Ca (bidentate ligand).

Proteins known to contain EF-hand regions include: Calmodulin (Ca=4, except in yeast where Ca=3) ("Ca=" indicates approximate number of EF-hand regions); diacylglycerol kinase (EC 2.7.1.107) (DGK) (Ca=2); 2) FAD-dependent glycerol-3-phosphate dehydrogenase (EC 1.1.99.5) from mammals (Ca=1); guanylate cyclase activating protein (GCAP) (Ca=3); MIF related proteins 8 (MRP-8 or CFAG) and 14 (MRP-14) (Ca=2); myosin regulatory light chains (Ca=1); oncomodulin (Ca=2); osteonectin (basement membrane protein BM-40) (SPARC); and proteins that contain an "osteonectin" domain (QR1, matrix glycoprotein SC1).

The consensus pattern includes the complete EF-hand loop as well as the first residue which follows the loop and which seem to always be hydrophobic: D-x-[DNS]-{ILVFYW}-[DENSTG]-[DNQGHRK]-{GP}-[LIVMC]-[DENQSTAGC]-x(2)-[DE]-

n) Ets Domain (Ets Nterm). SEQ ID NO:2849, and thus the sequence it validates, represents a polynucleotide encoding a polypeptide with N-terminal homology in ETS domain. Proteins of this family contain a conserved domain, the "ETS-domain," that is involved in DNA binding. The domain appears to recognize purine-rich sequences; it is about 85 to 90 amino acids in length, and is rich in aromatic and positively charged residues (Wasylyk, et al., , Eur. J. Biochem. (1993) 211:718).

The *ets* gene family encodes a novel class of DNA-binding proteins, each of which binds a specific DNA sequence. These proteins comprise an *ets* domain that specifically interacts with sequences containing the common core tri-nucleotide sequence GGA. In addition to an *ets* domain, native *ets* proteins comprise other sequences which can modulate the biological specificity of the protein. *Ets* genes and proteins are involved in a variety of essential biological processes including cell growth, differentiation and development, and three members are implicated in oncogenic process.

o) Type II fibronectin collagen-binding domain (FntypeII). A few of the validation sequences, and thus the sequences they validate, represent polynucleotides encoding a polypeptide having a type II fibronectin collagen binding domain. Fibronectin is a plasma protein that binds cell surfaces and various compounds including collagen, fibrin, heparin,

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DNA, and actin. The major part of the sequence of fibronectin consists of the repetition of three types of domains, which are called type I, II, and III (Skorstengaard K., et al., Eur. J. Biochem. (1986) 161:441). Type II domain is approximately forty residues long, contains four conserved cysteines involved in disulfide bonds and is part of the collagen-binding region of fibronectin. In fibronectin the type II domain is duplicated. Type II domains 5 have also been found in the following proteins: 1) blood coagulation factor XII (Hageman factor) (1 copy); 2) bovine seminal plasma proteins PDC-109 (BSP-A1/A2) and BSP-A3 (Seidah N.G., et al., Biochem. J. (1987) 243:195. (twice); 3) cation-independent mannose-6-phosphate receptor (which is also the insulin-like growth factor II receptor) Kornfeld S., Annu. Rev. Biochem. (1992) 61:307) (1 copy); 4) Mannose receptor of macrophages 10 (Taylor M.E., et al., J. Biol. Chem. (1990) 265:12156) (1 copy); 5) 180 Kd secretory phospholipase A2 receptor (1 copy) Lambeau G., et al., J. Biol. Chem. (1994) 269:1575; 6) DEC-205 receptor (1 copy); 6) Jiang W., et al., Nature (1995) 375:151); 7) 72 Kd type IV collagenase (EC 3.4.24.24) (MMP-2) (Collier I.E., et al., J. Biol. Chem. (1988) 263:6579) (3 copies); 7) 92 Kd type IV collagenase (EC 3.4.24.24) (MMP-9) (3 copies); 8) 15 Hepatocyte growth factor activator (Miyazawa K., et al., J. Biol. Chem. (1993) 268:10024) (1 copy).

A schematic representation of the position of the invariant residues and the topology of the disulfide bonds in fibronectin type II domain is shown below:

xxCxxPFx#xxxxxxCxxxxxxxWCxxxxx#xxx#x#Cxx where 'C' represents the conserved cysteine involved in a disulfide bond and '#' represents a large hydrophobic residue. The consensus pattern for identifying members of this family, which pattern spans this entire domain, is: C-x(2)-P-F-x-[FYWI]-x(7)-C-x(8,10)-W-C-x(4)-[DNSR]-[FYW]-x(3,5)-[FYW]-x-[FYWI]-C (where the four C's are involved in disulfide bonds).

p) G-Protein Alpha Subunit (G-alpha). Several of the validation sequences, and thus the sequences they validate, correspond to a gene encoding a novel polypeptide of the G-protein alpha subunit family. Guanine nucleotide binding proteins (G-proteins) are a family of membrane-associated proteins that couple extracellularly-activated integralmembrane receptors to intracellular effectors, such as ion channels and enzymes that vary the concentration of second messenger molecules. G-proteins are composed of 3 subunits (alpha, beta and gamma) which, in the resting state, associate as a trimer at the inner face of

the plasma membrane. The alpha subunit has a molecule of guanosine diphosphate (GDP) bound to it. Stimulation of the G-protein by an activated receptor leads to its exchange for GTP (guanosine triphosphate). This results in the separation of the alpha from the beta and gamma subunits, which always remain tightly associated as a dimer. Both the alpha and beta-gamma subunits are then able to interact with effectors, either individually or in a cooperative manner. The intrinsic GTPase activity of the alpha subunit hydrolyses the bound GTP to GDP. This returns the alpha subunit to its inactive conformation and allows it to reassociate with the beta-gamma subunit, thus restoring the system to its resting state.

G-protein alpha subunits are 350-400 amino acids in length and have molecular weights in the range 40-45 kDa. Seventeen distinct types of alpha subunit have been identified in mammals. These fall into 4 main groups on the basis of both sequence similarity and function: alpha-s, alpha-q, alpha-i and alpha-12 (Simon *et al.*, *Science* (1993) 252:802). Many alpha subunits are substrates for ADP-ribosylation by cholera or pertussis toxins. They are often N-terminally acylated, usually with myristate and/or palmitoylate, and these fatty acid modifications are probably important for membrane association and high- affinity interactions with other proteins. The atomic structure of the alpha subunit of the G-protein involved in mammalian vision, transducin, has been elucidated in both GTP- and GDB-bound forms, and shows considerable similarity in both primary and tertiary structure in the nucleotide-binding regions to other guanine nucleotide binding proteins, such as p21-ras and EF-Tu.

- q) <u>Helicases conserved C-terminal domain (helicase C)</u>. SEQ ID NOS:2503, 4469, and 5020, and thus the sequences they validate, represent polynucleotides encoding novel members of the DEAD/H helicase family. The DEAD and DEAH families are described above.
- r) Homeobox domain (homeobox). SEQ ID NO:4241, and thus the sequence it validates, represents a polynucleotide encoding a protein having a homeobox domain. The 'homeobox' is a protein domain of 60 amino acids (Gehring In: Guidebook to the Homebox Genes, Duboule D., Ed., pp1-10, Oxford University Press, Oxford, (1994); Buerglin In: Guidebook to the Homebox Genes, pp25-72, Oxford University Press, Oxford, (1994); Gehring Trends Biochem. Sci. (1992) 17:277-280; Gehring et alAnnu. Rev. Genet. (1986) 20:147-173; Schofield Trends Neurosci. (1987) 10:3-6; http://copan.bioz.unibas.ch/homeo.html) first identified in number of Drosophila homeotic and segmentation proteins. It is extremely well conserved in many other animals, including vertebrates. This domain

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binds DNA through a helix-turn-helix type of structure. Several proteins that contain a homeobox domain play an important role in development. Most of these proteins are sequence-specific DNA-binding transcription factors. The homeobox domain is also very similar to a region of the yeast mating type proteins. These are sequence-specific DNA-binding proteins that act as master switches in yeast differentiation by controlling gene expression in a cell type-specific fashion.

A schematic representation of the homeobox domain is shown below. The helix-turn-helix region is shown by the symbols 'H' (for helix), and 't' (for turn).

The pattern detects homeobox sequences 24 residues long and spans positions 34 to 57 of the homeobox domain. The consensus pattern is as follows: [LIVMFYG]-[ASLVR]-x(2)-[LIVMSTACN]-x-[LIVM]-x(4)-[LIV]-[RKNQESTAIY]-[LIVFSTNKH]-W-[FYVC]-x-[NDQTAH]-x(5)-[RKNAIMW].

x) MAP kinase kinase (mkk). Several validation sequences, and thus the sequences they validate, represent novel members of the MAP kinase kinase family. MAP kinases (MAPK) are involved in signal transduction, and are important in cell cycle and cell growth controls. The MAP kinase kinases (MAPKK) are dual-specificity protein kinases which phosphorylate and activate MAP kinases. MAPKK homologues have been found in yeast, invertebrates, amphibians, and mammals. Moreover, the MAPKK/MAPK phosphorylation switch constitutes a basic module activated in distinct pathways in yeast and in vertebrates. MAPKK regulation studies have led to the discovery of at least four MAPKK convergent pathways in higher organisms. One of these is similar to the yeast pheromone response pathway which includes the stell protein kinase. Two other pathways require the activation of either one or both of the serine/threonine kinase-encoded oncogenes c-Raf-1 and c-Mos. Additionally, several studies suggest a possible effect of the cell cycle control regulator cyclin-dependent kinase 1 (cdc2) on MAPKK activity. Finally, MAPKKs are apparently essential transducers through which signals must pass

Finally, MAPKKs are apparently essential transducers through which signals must pass before reaching the nucleus. For review, see, e.g., Biologique Biol Cell (1993) 79:193-207; Nishida et al., Trends Biochem Sci (1993) 18:128-31; Ruderman Curr Opin Cell Biol (1993) 5:207-13; Dhanasekaran et al., Oncogene (1998) 17:1447-55; Kiefer et al., Biochem Soc Trans (1997) 25:491-8; and Hill, Cell Signal (1996) 8:533-44.

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y) 3'5'-cyclic nucleotide phosphodiesterases signature (PDEase). SEQ ID NO:4482, and thus the sequence it validates, represents a polynucleotide encoding a novel 3'5'-cyclic nucleotide phosphodiesterases (PDEases). PDEases catalyze the hydrolysis of cAMP or cGMP to the corresponding nucleoside 5' monophosphates (Charbonneau H., et al, *Proc. Natl. Acad. Sci. U.S.A.* (1986) 83:9308). There are at least seven different subfamilies of PDEases (Beavo J.A., et al., *Trends Pharmacol. Sci.* (1990) 11:150; http://weber.u.washington.edu/~pde/: 1) Type 1, calmodulin/calcium-dependent PDEases; 2) Type 2, cGMP-stimulated PDEases; 3) Type 3, cGMP-inhibited PDEases; 4) Type 4, cAMP-specific PDEases.; 5) Type 5, cGMP-specific PDEases; 6) Type 6, rhodopsin-sensitive cGMP-specific PDEases; and 7) Type 7, High affinity cAMP-specific PDEases.

All PDEase forms share a conserved domain of about 270 residues. The signature pattern is determined from a stretch of 12 residues that contains two conserved histidines: H-D-[LIVMFY]-x-H-x-[AG]-x(2)-[NQ]-x-[LIVMFY].

z) Protein Kinase (protkinase). Several validation sequences, and thus the sequences they validate, represent polynucleotides encoding protein kinases. Protein kinases catalyze phosphorylation of proteins in a variety of pathways, and are implicated in cancer. Eukaryotic protein kinases (Hanks S.K., et al., FASEB J. (1995) 9:576; Hunter T., Meth. Enzymol. (1991) 200:3; Hanks S.K., et al., Meth. Enzymol. (1991) 200:38; Hanks S.K., Curr. Opin. Struct. Biol. (1991) 1:369; Hanks S.K., et al., Science (1988) 241:42) are enzymes that belong to a very extensive family of proteins which share a conserved catalytic core common to both serine/threonine and tyrosine protein kinases. There are a number of conserved regions in the catalytic domain of protein kinases. Two of the conserved regions are the basis for the signature pattern in the protein kinase profile. The first region, which is located in the N-terminal extremity of the catalytic domain, is a glycine-rich stretch of residues in the vicinity of a lysine residue, which has been shown to be involved in ATP binding. The second region, which is located in the central part of the catalytic domain, contains a conserved aspartic acid residue which is important for the catalytic activity of the enzyme (Knighton D.R., et al., Science (1991) 253:407). The protein kinase profile includes two signature patterns for this second region: one specific for serine/threonine kinases and the other for tyrosine kinases. A third profile is based on the alignment in (Hanks S.K., et al., FASEB J. (1995) 9:576) and covers the entire catalytic domain. The consensus patterns are as follows:

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- 1) Consensus pattern: [LIV]-G-{P}-G-{P}-[FYWMGSTNH]-[SGA]-{PW}-[LIVCAT]-{PD}-x-[GSTACLIVMFY]-x(5,18)-[LIVMFYWCSTAR]-[AIVP]-[LIVMFAGCKR]-K, where K binds ATP. The majority of known protein kinases are detected by this pattern. Proteins kinases that are not detected by this consensus include viral kinases, which are quite divergent in this region and are completely missed bythis pattern.
- 2) Consensus pattern: [LIVMFYC]-x-[HY]-x-D-[LIVMFY]-K-x(2)-N-[LIVMFYCT](3), where D is an active site residue. This consensus sequence identifies most serine/threonine-specific protein kinases with only 10 exceptions. Half of the exceptions are viral kinases, while the other exceptions include Epstein-Barr virus BGLF4 and Drosophila ninaC, which have Ser and Arg, respectively, instead of the conserved Lys. These latter two protein kinases are detected by the tyrosine kinase specific pattern described below.
- 3) Consensus pattern: [LIVMFYC]-x-[HY]-x-D-[LIVMFY]-[RSTAC]-x(2)-N-[LIVMFYC], where D is an active site residue. All tyrosine-specific protein kinases are detected by this consensus pattern, with the exception of human ERBB3 and mouse blk. This pattern also detects most bacterial aminoglycoside phosphotransferases (Benner S., *Nature* (1987) 329:21; Kirby R., *J. Mol. Evol.* (1992) 30:489) and herpesviruses ganciclovir kinases (Littler E., *et al.*, *Nature* (1992) 358:160), which are structurally and evolutionary related to protein kinases.

The protein kinase profile also detects receptor guanylate cyclases and 2-5A-dependent ribonucleases. Sequence similarities between these two families and the eukaryotic protein kinase family have been noticed previously. The profile also detects Arabidopsis thaliana kinase-like protein TMKL1 which seems to have lost its catalytic activity.

If a protein analyzed includes the two of the above protein kinase signatures, the probability of it being a protein kinase is close to 100%. Eukaryotic-type protein kinases have also been found in prokaryotes such as Myxococcus xanthus (Munoz-Dorado J., et al., Cell (1991) 67:995) and Yersinia pseudotuberculosis. The patterns shown above has been updated since their publication in (Bairoch A., et al., Nature (1988) 331:22).

aa) Ras family proteins (ras). SEQ IDNO:3671, and thus the sequence it validates, represent polynucleotides encoding the ras family of small GTP/GDP-binding proteins (Valencia et al., 1991, Biochemistry 30:4637-4648). Ras family members generally require

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a specific guanine nucleotide exchange factor (GEF) and a specific GTPase activating protein (GAP) as stimulators of overall GTPase activity. Among ras-related proteins, the highest degree of sequence conservation is found in four regions that are directly involved in guanine nucleotide binding. The first two constitute most of the phosphate and Mg2+ binding site (PM site) and are located in the first half of the G-domain. The other two regions are involved in guanosine binding and are located in the C-terminal half of the molecule. Motifs and conserved structural features of the ras-related proteins are described in Valencia et al., 1991, Biochemistry 30:4637-4648.

A major consensus pattern of ras proteins is: D-T-A-G-Q-E-K-[LF]-G-G-L-R-[DE]-G-Y-Y.

bb) Thioredoxin family active site (Thioredox). SEQ ID NO:3936, and thus the sequence it validates, represent a polynucleotide encoding a protein having a thioredoxin family active site. Thioredoxins (Holmgren A., Annu. Rev. Biochem. (1985) 54:237; Gleason F.K., et al., FEMS Microbiol. Rev. (1988) 54:271; Holmgren A. J. Biol. Chem. (1989) 264:13963; Eklund H., et al. Proteins (1991) 11:13) are small proteins of approximately one hundred amino- acid residues which participate in various redox reactions via the reversible oxidation of an active center disulfide bond. They exist in either a reduced form or an oxidized form where the two cysteine residues are linked in an intramolecular disulfide bond. Thioredoxin is present in prokaryotes and eukaryotes and the sequence around the redox-active disulfide bond is well conserved.

A number of eukaryotic proteins contain domains evolutionary related to thioredoxin, and all of them are protein disulphide isomerases (PDI). PDI (Freedman R.B., et al., *Biochem. Soc. Trans.* (1988) 16:96; Kivirikko K.I., et al., *FASEB J.* (1989) 3:1609; Freedman R.B., et al. *Trends Biochem. Sci.* (1994) 19:331) is an endoplasmic reticulum enzyme that catalyzes the rearrangement of disulfide bonds in various proteins. The various forms of PDI which are currently known are: 1) PDI major isozyme; a multifunctional protein that also function as the beta subunit of prolyl 4-hydroxylase (EC 1.14.11.2), as a component of oligosaccharyl transferase (EC 2.4.1.119), as thyroxine deiodinase, as glutathione-insulin transhydrogenase, and as a thyroid hormone-binding protein; 2) ERp60 (ER-60; 58 Kd microsomal protein), which is a protease; 3) ERp72; and 4) P5.

All PDI contains two or three (ERp72) copies of the thioredoxin domain. The consensus pattern is: [LIVMF]-[LIVMSTA]-x-[LIVMFYC]-[FYWSTHE]-x(2)-

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virus protein A53 (SalF19R).

[FYWGTN]-C- [GATPLVE]-[PHYWSTA]-C-x(6)-[LIVMFYWT] (where the two C's form the redox-active bond.

cc) TNFR/NGFR family cysteine-rich region (TNFR_c6). SEQ ID NO:3927, and thus the sequence it validates, represent a polynucleotide encoding a protein having a TNFR/NGFR family cysteine-rich region. A number of proteins, some of which are known to be receptors for growth factors, have been found to contain a cysteine-rich domain of about 110 to 160 amino acids in their N-terminal part, that can be subdivided into four (or in some cases, three) modules of about 40 residues containing 6 conserved cysteines. Proteins known to belong to this family (Mallet S., et al., Immunol. Today (1991) 12:220; Sprang S.R., Trends Biochem. Sci. (1990) 15:366; Krammer P.H., et al., Curr. Biol. (1992) 2:383; Bazan J.F., Curr. Biol. (1993) 3:603) are: 1) Tumor Necrosis Factor type I and type II receptors (TNFR) (Both receptors bind TNF-alpha and TNF-beta, but are only similar in the cysteine-rich region.); 2) Shope fibroma virus soluble TNF receptor (protein T2); 3) Lymphotoxin alpha/beta receptor; 4) Low-affinity nerve growth factor receptor (LA-NGFR); 5) CD40 (Bp50), the receptor for the CD40L (or TRAP) cytokine; 6) CD27, the receptor for the CD27L cytokine; 8) CD30, the receptor for the CD30L cytokine; 9) T-cell protein 4-1BB, the receptor for the 4-1BBL putative cytokine; 10) FAS antigen (or APO-1), the receptor for FASL, a protein involved in apoptosis (programmed cell death); 11) T-cell antigen OX40, the receptor for the OX40L cytokine; 12) Wsl-1, a receptor (for a yet undefined ligand) that mediates apoptosis; 13) Vaccinia

The six cysteines all involved in intrachain disulfide bonds (Banner D.W., et al, Cell (1993) 73:431). A schematic representation of the structure of the 40 residue module of these receptors is shown below:

- - dd) <u>Four Transmembrane Integral Membrane Proteins (transmembrane4)</u>. Several of the validation sequences, and thus the sequences they validate, correspond to a sequence encoding a polypeptide that is a member of the 4 transmembrane segments integral

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membrane protein family (transmembrane 4 family). The transmembrane 4 family of proteins includes a number of evolutionarily-related eukaryotic cell surface antigens (Levy et al., J. Biol. Chem., (1991) 266:14597; Tomlinson et al., Eur. J. Immunol. (1993) 23:136; Barclay et al. The leucocyte antigen factbooks. (1993) Academic Press, London/San

- Diego). The proteins belonging to this family include: 1) Mammalian antigen CD9 (MIC3), which is involved in platelet activation and aggregation; 2) Mammalian leukocyte antigen CD37, expressed on B lymphocytes; 3) Mammalian leukocyte antigen CD53 (OX-44), which is implicated in growth regulation in hematopoietic cells; 4) Mammalian lysosomal membrane protein CD63 (melanoma-associated antigen ME491; antigen AD1);
- 5) Mammalian antigen CD81 (cell surface protein TAPA-1), which is implicated in regulation of lymphoma cell growth; 6) Mammalian antigen CD82 (protein R2; antigen C33; Kangai 1 (KAI1)), which associates with CD4 or CD8 and delivers costimulatory signals for the TCR/CD3 pathway; 7) Mammalian antigen CD151 (SFA-1; platelet-endothelial tetraspan antigen 3 (PETA-3)); 8) Mammalian cell surface glycoprotein A15
 (TALLA-1; MXS1); 9) Mammalian novel antigen 2 (NAG-2); 10) Human tumor-associated antigen CO-029; 11) Schistosoma mansoni and japonicum 23 Kd surface antigen (SM23 / SJ23).

The members of the 4 transmembrane family share several characteristics. First, they all are apparently type III membrane proteins, which are integral membrane proteins containing an N-terminal membrane-anchoring domain which is not cleaved during biosynthesis and which functions both as a translocation signal and as a membrane anchor. The family members also contain three additional transmembrane regions, at least seven conserved cysteines residues, and are of approximately the same size (218 to 284 residues). These proteins are collectively know as the "transmembrane 4 superfamily" (TM4) because they span plasma membrane four times.

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A schematic diagram of the domain structure of these proteins is as follows:

where Cyt is the cytoplasmic domain, TMa is the transmembrane anchor; TM2 to TM4 represents transmembrane regions 2 to 4, 'C' are conserved cysteines, and '*'indicates the position of the consensus pattern. The consensus pattern spans a conserved region including two cysteines located in a short cytoplasmic loop between two transmembrane domains: Consensus pattern: G-x(3)-[LIVMF]-x(2)-[GSA]-[LIVMF](2)-G-C-x-[GA]-[STA]- x(2)-[EG]-x(2)-[CWN]-[LIVM](2).

ee) Trypsin (trypsin). SEQ ID NOS:3381, 4684, and 4688, and thus the sequences they validate, correspond to novel serine proteases of the trypsin family. The catalytic activity of the serine proteases from the trypsin family is provided by a charge relay system involving an aspartic acid residue hydrogen-bonded to a histidine, which itself is hydrogenbonded to a serine. The sequences in the vicinity of the active site serine and histidine residues are well conserved in this family of proteases (Brenner S., Nature (1988) 334:528). Proteases known to belong to the trypsin family include: 1) Acrosin; 2) Blood coagulation factors VII, IX, X, XI and XII, thrombin, plasminogen, and protein C; 3) Cathepsin G; 4) Chymotrypsins; 5) Complement components C1r, C1s, C2, and complement factors B, D and I; 6) Complement-activating component of RA-reactive factor; 7) Cytotoxic cell proteases (granzymes A to H); 8) Duodenase I; 9) Elastases 1, 2, 3A, 3B (protease E), leukocyte (medullasin).; 10) Enterokinase (EC 3.4.21.9) (enteropeptidase); 11) Hepatocyte growth factor activator; 12) Hepsin; 13) Glandular (tissue) kallikreins (including EGF-binding protein types A, B, and C, NGF-gamma chain, gamma-renin, prostate specific antigen (PSA) and tonin); 14) Plasma kallikrein; 15) Mast cell proteases (MCP) 1 (chymase) to 8; 16) Myeloblastin (proteinase 3) (Wegener's autoantigen); 17) Plasminogen activators (urokinase-type, and tissue-type); 18) Trypsins I, II, III, and IV; 19) Tryptases; 20) Snake venom proteases such as ancrod, batroxobin, cerastobin, flavoxobin, and protein C activator; 21) Collagenase from common cattle grub and collagenolytic protease from Atlantic sand fiddler crab; 22) Apolipoprotein(a); 23) Blood fluke cercarial protease; 24) Drosophila trypsin like proteases: alpha, easter, snakelocus; 25) Drosophila protease stubble (gene sb); and 26) Major mite fecal allergen Der p

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III. All the above proteins belong to family S1 in the classification of peptidases (Rawlings N.D., et al., Meth. Enzymol. (1994) 244:19; http://www.expasy.ch/cgibin/lists?peptidas.txt) and originate from eukaryotic species. It should be noted that bacterial proteases that belong to family S2A are similar enough in the regions of the active site residues that they can be picked up by the same patterns.

The consensus patterns for this trypsin protein family are: 1) [LIVM]-[ST]-A-[STAG]-H-C, where H is the active site residue. All sequences known to belong to this class detected by the pattern, except for complement components C1r and C1s, pig plasminogen, bovine protein C, rodent urokinase, ancrod, gyroxin and two insect trypsins; 2) [DNSTAGC]-[GSTAPIMVQH]-x(2)-G-[DE]-S-G-[GS]-[SAPHV]- [LIVMFYWH]-[LIVMFYSTANQH], where S is the active site residue. All sequences known to belong to this family are detected by the above consensus sequences, except for 18 different proteases which have lost the first conserved glycine. If a protein includes both the serine and the histidine active site signatures, the probability of it being a trypsin family serine protease is 100%.

ff) <u>WD Domain, G-Beta Repeats (WD domain)</u>. A few of the validation sequences, and the sequences they validate, represent novel members of the WD domain/G-beta repeat family. Beta-transducin (G-beta) is one of the three subunits (alpha, beta, and gamma) of the guanine nucleotide-binding proteins (G proteins) which act as intermediaries in the transduction of signals generated by transmembrane receptors (Gilman, *Annu. Rev. Biochem.* (1987) 56:615). The alpha subunit binds to and hydrolyzes GTP; the functions of the beta and gamma subunits are less clear but they seem to be required for the replacement of GDP by GTP as well as for membrane anchoring and receptor recognition.

In higher eukaryotes, G-beta exists as a small multigene family of highly conserved proteins of about 340 amino acid residues. Structurally, G-beta consists of eight tandem repeats of about 40 residues, each containing a central Trp-Asp motif (this type of repeat is sometimes called a WD-40 repeat). Such a repetitive segment has been shown to exist in a number of other proteins including: human LIS1, a neuronal protein involved in type-1 lissencephaly; and mammalian coatomer beta' subunit (beta'-COP), a component of a cytosolic protein complex that reversibly associates with Golgi membranes to form vesicles that mediate biosynthetic protein transport.

The consensus pattern for the WD domain/G-Beta repeat family is: [LIVMSTAC]-

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[LIVMFYWSTAGC]-[LIMSTAG]-[LIVMSTAGC]-x(2)-[DN]-x(2)-[LIVMWSTAC]-x-[LIVMFSTAG]-W-[DEN]-[LIVMFSTAGCN].

gg) wnt Family of Developmental Signaling Proteins (Wnt_dev_sign). Several of the validation sequences, and thus the sequences they validate, correspond to novel members of the wnt family of developmental signaling proteins. Wnt-1 (previously known as int-1), the seminal member of this family, (Nusse R., *Trends Genet.* (1988) 4:291) is a proto-oncogene induced by the integration of the mouse mammary tumor virus. It is thought to play a role in intercellular communication and seems to be a signalling molecule important in the development of the central nervous system (CNS). The sequence of wnt-1 is highly conserved in mammals, fish, and amphibians. Wnt-1 was found to be a member of a large family of related proteins (Nusse R., *et al.*, *Cell* (1992) 69:1073; McMahon A.P., *Trends Genet.* (1992) 8:1; Moon R.T., *BioEssays* (1993) 15:91) that are all thought to be developmental regulators. These proteins are known as wnt-2 (also known as irp), wnt-3, -3A, -4, -5A, -5B, -6, -7A, -7B, -8, -8B, -9 and -10. At least four members of this family are present in Drosophila; one of them, wingless (wg), is implicated in segmentation polarity.

All these proteins share the following features characteristics of secretory proteins: a signal peptide, several potential N-glycosylation sites and 22 conserved cysteines that are probably involved in disulfide bonds. The Wnt proteins seem to adhere to the plasma membrane of the secreting cells and are therefore likely to signal over only few cell diameters. The consensus pattern, which is based upon a highly conserved region including three cysteines, is as follows: C-K-C-H-G-[LIVMT]-S-G-x-C. All sequences known to belong to this family are detected by the provided consensus pattern.

hh) Protein Tyrosine Phosphatase (Y_phosphatase). Several of the validation sequences, and thus the sequences they validate, represent a polynucleotide encoding a protein tyrosine kinase. Tyrosine specific protein phosphatases (EC 3.1.3.48) (PTPase) (Fischer et al., Science (1991) 253:401; Charbonneau et al., Annu. Rev. Cell Biol. (1992) 8:463; Trowbridge, J. Biol. Chem. (1991) 266:23517; Tonks et al., Trends Biochem. Sci. (1989) 14:497; and Hunter, Cell (1989) 58:1013) catalyze the removal of a phosphate group attached to a tyrosine residue. These enzymes are very important in the control of cell growth, proliferation, differentiation and transformation. Multiple forms of PTPase have been characterized and can be classified into two categories: soluble PTPases and transmembrane receptor proteins that contain PTPase domain(s).

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Soluble PTPases include PTPN3 (H1) and PTPN4 (MEG), enzymes that contain an N-terminal band 4.1-like domain and could act at junctions between the membrane and cytoskeleton; PTPN6 (PTP-1C; HCP; SHP) and PTPN11 (PTP-2C; SH-PTP3; Syp), enzymes that contain two copies of the SH2 domain at its N-terminal extremity.

Dual specificity PTPases include DUSP1 (PTPN10; MAP kinase phosphatase-1; MKP-1) which dephosphorylates MAP kinase on both Thr-183 and Tyr-185; and DUSP2 (PAC-1), a nuclear enzyme that dephosphorylates MAP kinases ERK1 and ERK2 on both Thr and Tyr residues.

Structurally, all known receptor PTPases are made up of a variable length extracellular domain, followed by a transmembrane region and a C-terminal catalytic cytoplasmic domain. Some of the receptor PTPases contain fibronectin type III (FN-III) repeats, immunoglobulin-like domains, MAM domains or carbonic anhydrase-like domains in their extracellular region. The cytoplasmic region generally contains two copies of the PTPAse domain. The first seems to have enzymatic activity, while the second is inactive but seems to affect substrate specificity of the first. In these domains, the catalytic cysteine is generally conserved but some other, presumably important, residues are not.

PTPase domains consist of about 300 amino acids. There are two conserved cysteines and the second one has been shown to be absolutely required for activity. Furthermore, a number of conserved residues in its immediate vicinity have also been shown to be important. The consensus pattern for PTPases is: [LIVMF]-H-C-x(2)-G-x(3)-[STC]-[STAGP]-x-[LIVMFY]; C is the active site residue.

<u>ii)Zinc Finger, C2H2 Type (Zincfing_C2H2)</u>. Several of the validation sequences, and thus the sequences they validate, correspond to polynucleotides encoding novel members of the of the C2H2 type zinc finger protein family. Zinc finger domains (Klug et al., Trends Biochem. Sci. (1987) 12:464; Evans et al., Cell (1988) 52:1; Payre et al., FEBS Lett. (1988) 234:245; Miller et al., EMBO J. (1985) 4:1609; and Berg, Proc. Natl. Acad. Sci. USA (1988) 85:99) are nucleic acid-binding protein structures first identified in the Xenopus transcription factor TFIIIA. These domains have since been found in numerous nucleic acid-binding proteins. A zinc finger domain is composed of 25 to 30 amino acid residues. Two cysteine or histidine residues are positioned at both extremities of the domain, which are involved in the tetrahedral coordination of a zinc atom. It has been proposed that such a domain interacts with about five nucleotides.

Many classes of zinc fingers are characterized according to the number and

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positions of the histidine and cysteine residues involved in the zinc atom coordination. In the first class to be characterized, called C2H2, the first pair of zinc coordinating residues are cysteines, while the second pair are histidines. A number of experimental reports have demonstrated the zinc-dependent DNA or RNA binding property of some members of this class.

Mammalian proteins having a C2H2 zipper include (number in parenthesis indicates number of zinc finger regions in the protein): basonuclin (6), BCL-6/LAZ-3 (6), erythroid krueppel-like transcription factor (3), transcription factors Sp1 (3), Sp2 (3), Sp3 (3) and Sp(4) 3, transcriptional repressor YY1 (4), Wilms' tumor protein (4), EGR1/Krox24 (3), EGR2/Krox20 (3), EGR3/Pilot (3), EGR4/AT133 (4), Evi-1 (10), GLI1 (5), GLI2 (4+), GLI3 (3+), HIV-EP1/ZNF40 (4), HIV-EP2 (2), KR1 (9+), KR2 (9), KR3 (15+), KR4 (14+), KR5 (11+), HF.12 (6+), REX-1 (4), ZfX (13), ZfY (13), Zfp-35 (18), ZNF7 (15), ZNF8 (7), ZNF35 (10), ZNF42/MZF-1 (13), ZNF43 (22), ZNF46/Kup (2), ZNF76 (7), ZNF91 (36), ZNF133 (3).

In addition to the conserved zinc ligand residues, it has been shown that a number of other positions are also important for the structural integrity of the C2H2 zinc fingers. (Rosenfeld *et al.*, *J. Biomol. Struct. Dyn.* (1993) 11:557) The best conserved position is found four residues after the second cysteine; it is generally an aromatic or aliphatic residue. The consensus pattern for C2H2 zinc fingers is: C-x(2,4)-C-x(3)-[LIVMFYWC]-x(8)-H-x(3,5)-H. The two C's and two H's are zinc ligands.

- jj) Zinc finger, C3HC4 type (RING finger), signature (Zincfing C3H4). SEQ ID NOS:3774 and 4477, and thus the sequences they validate, represent polynucleotides encoding a polypeptide having a C3HC4 type zinc finger signature. A number of eukaryotic and viral proteins contain this signature, which is primarily a conserved cysteine-rich domain of 40 to 60 residues (Borden K.L.B., et al., *Curr. Opin. Struct. Biol.* (1996) 6:395) that binds two atoms of zinc, and is probably involved in mediating protein-protein interactions. The 3D structure of the zinc ligation system is unique to the RING domain and is referred to as the "cross-brace" motif. The spacing of the cysteines in such a domain is C-x(2)-C-x(9 to 39)-C-x(1 to 3)-H-x(2 to 3)-C-x(2)-C-x(4 to 48)-C-x(2)-C. Proteins that include the C3HC4 domain include:
- 1) Mammalian V(D)J recombination activating protein (RAG1). RAG1 activates the rearrangement of immunoglobulin and T-cell receptor genes.
 - 2) Mouse rpt-1. Rpt-1 is a trans-acting factor that regulates gene expression directed

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by the promoter region of the interleukin-2 receptor alpha chainor the LTR promoter region of HIV-1.

- 3) Human rfp. Rfp is a developmentally regulated protein that may function in male germ cell development. Recombination of the N-terminal section of rfp with a protein tyrosine kinase produces the ret transforming protein.
- 4) Human 52 Kd Ro/SS-A protein. A protein of unknown function from the Ro/SS-A ribonucleoprotein complex. Sera from patients with systemic lupus erythematosus or primary Sjogren's syndrome often contain antibodies that react with the Ro proteins.
 - 5) Human histocompatibility locus protein RING1.
- 6) Human PML, a probable transcription factor. Chromosomal translocation of PML with retinoic receptor alpha creates a fusion protein which is the cause of acute promyelocytic leukemia (APL).
- 7) Mammalian breast cancer type 1 susceptibility protein (BRCA1) ([E1] http://bioinformatics.weizmann.ac.il/hotmolecbase/entries/brca1.htm).
 - 8) Mammalian cbl proto-oncogene.
 - 9) Mammalian bmi-1 proto-oncogene.
- 10) Vertebrate CDK-activating kinase (CAK) assembly factor MAT1, a protein that stabilizes the complex between the CDK7 kinase and cyclin H (MAT1 stands for 'Menage A Trois').
- 11) Mammalian mel-18 protein. Mel-18 which is expressed in a variety of tumorcells is a transcriptional repressor that recognizes and bind a specific DNA sequence.
- 12) Mammalian peroxisome assembly factor-1 (PAF-1) (PMP35), which is somewhat involved in the biogenesis of peroxisomes. In humans, defects in PAF-1 are responsible for a form of Zellweger syndrome, an autosomal recessive disorder associated with peroxisomal deficiencies.
 - 13) Human MAT1 protein, which interacts with the CDK7-cyclin H complex.
 - 14) Human RING1 protein.
 - 15) Xenopus XNF7 protein, a probable transcription factor.
- 16) Trypanosoma protein ESAG-8 (T-LR), which may be involved in the postranscriptional regulation of genes in VSG expression sites or may interact with adenylate cyclase to regulate its activity.
 - 17) Drosophila proteins Posterior Sex Combs (Psc) and Suppressor two of zeste

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- (Su(z)2). The two proteins belong to the Polycomb group of genes needed to maintain the segment-specific repression of homeotic selector genes.
- 18) Drosophila protein male-specific msl-2, a DNA-binding protein which is involved in X chromosome dosage compensation (the elevation of transcription of the male single X chromosome).
- 19) Arabidopsis thaliana protein COP1 which is involved in the regulation ofphotomorphogenesis.
 - 20) Fungal DNA repair proteins RAD5, RAD16, RAD18 and rad8.
- 21) Herpesviruses trans-acting transcriptional protein ICP0/IE110. This protein which has been characterized in many different herpesviruses is a trans-activator and/or repressor of the expression of many viral and cellular promoters.
 - 22) Baculoviruses protein CG30.
 - 23) Baculoviruses major immediate early protein (PE-38).
 - 24) Baculoviruses immediate-early regulatory protein IE-N/IE-2.
 - 25) Caenorhabditis elegans hypothetical proteins F54G8.4, R05D3.4 and T02C1.1.
 - 26) Yeast hypothetical proteins YER116c and YKR017c.

The signature pattern for the C3HC4 finger is based on the central region of the domain:

Consensus pattern: C-x-H-x-[LIVMFY]-C-x(2)-C-[LIVMYA].

Example 4: Differential Expression of Polynucleotides of the Invention: Description of Libraries and Detection of Differential Expression

The relative expression levels of the polynucleotides of the invention was assessed in several libraries prepared from various sources, including cell lines and patient tissue samples. Table 4 provides a summary of these libraries, including the shortened library name (used hereafter), the mRNA source used to prepared the cDNA library, the "nickname" of the library that is used in the tables below (in quotes), and the approximate number of clones in the library.

30 Table 4 Description of cDNA Libraries

Library	Description	Number of
(lib #)		Ciones in this Clustering
1	Km12 L4	

Library	Description	Number of Clones in this
(lib #)		Clustering
	Human Colon Cell Line, High Metastatic Potential (derived from Km12C) "High Colon"	307133
2	Km12C Human Colon Cell Line, Low Metastatic Potential "Low Colon"	284755
3	MDA-MB-231 Human Breast Cancer Cell Line, High Metastatic Potential; micro- metastases in lung "High Breast"	326937
4	MCF7 Human Breast Cancer Cell, Non Metastatic "Low Breast"	318979
8	MV-522 Human Lung Cancer Cell Line, High Metastatic Potential "High Lung"	223620
9	UCP-3 Human Lung Cancer Cell Line, Low Metastatic Potential "Low Lung"	312503
12	Human microvascular endothelial cells (HMEC) – Untreated PCR (OligodT) cDNA library	41938
13	Human microvascular endothelial cells (HMEC) – Basic fibroblast growth factor (bFGF) treated PCR (OligodT) cDNA library	42100
14	Human microvascular endothelial cells (HMEC) – Vascular endothelial growth factor (VEGF) treated PCR (OligodT) cDNA library	42825
15	Normal Colon – UC#2 Patient PCR (OligodT) cDNA library "Normal Colon Tumor Tissue"	34285
16	Colon Tumor – UC#2 Patient PCR (OligodT) cDNA library "Normal Colon Tumor Tissue"	35625
17	Liver Metastasis from Colon Tumor of UC#2 Patient PCR (OligodT) cDNA library "High Colon Metastasis Tissue"	36984
18	Normal Colon – UC#3 Patient PCR (OligodT) cDNA library "Normal Colon Tumor Tissue"	36216
19	Colon Tumor – UC#3 Patient PCR (OligodT) cDNA library "High Colon Tumor Tissue"	41388
20	Liver Metastasis from Colon Tumor of UC#3 Patient PCR (OligodT) cDNA library "High Colon Metastasis Tissue"	30956

The KM12L4 and KM12C cell lines are described in Example 1 above. The MDA-MB-231 cell line was originally isolated from pleural effusions (Cailleau, *J. Natl. Cancer. Inst.* (1974) 53:661), is of high metastatic potential, and forms poorly differentiated

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adenocarcinoma-grade II in nude mice consistent with breast carcinoma. The MCF7 cell line was derived from a pleural effusion of a breast adenocarcinoma and is non-metastatic. The MV-522 cell line is derived from a human lung carcinoma and is of high metastatic potential. The UCP-3 cell line is a low metastatic human lung carcinoma cell line; the 5 MV-522 is a high metastatic variant of UCP-3. These cell lines are well-recognized in the art as models for the study of human breast and lung cancer (see, e.g., Chandrasekaran et al., Cancer Res. (1979) 39:870 (MDA-MB-231 and MCF-7); Gastpar et al., J Med Chem (1998) 41:4965 (MDA-MB-231 and MCF-7); Ranson et al., Br J Cancer (1998) 77:1586 (MDA-MB-231 and MCF-7); Kuang et al., Nucleic Acids Res (1998) 26:1116 (MDA-MB-10 231 and MCF-7); Varki et al., Int J Cancer (1987) 40:46 (UCP-3); Varki et al., Tumour Biol. (1990) 11:327; (MV-522 and UCP-3); Varki et al., Anticancer Res. (1990) 10:637; (MV-522); Kelner et al., Anticancer Res (1995) 15:867 (MV-522); and Zhang et al., Anticancer Drugs (1997) 8:696 (MV522)). The samples of libraries 15-20 are derived from two different patients (UC#2, and UC#3). The bFGF-treated HMEC were prepared 15 by incubation with bFGF at 10ng/ml for 2 hrs; the VEGF-treated HMEC were prepared by incubation with 20ng/ml BEGF for 2 hrs. Following incubation with the respective growth factor, the cells were washed and lysis buffer added for RNA preparation.

Each of the libraries is composed of a collection of cDNA clones that in turn are representative of the mRNAs expressed in the indicated mRNA source. In order to facilitate the analysis of the millions of sequences in each library, the sequences were assigned to clusters. The concept of "cluster of clones" is derived from a sorting/grouping of cDNA clones based on their hybridization pattern to a panel of roughly 300 7bp oligonucleotide probes (see Drmanac et al., Genomics (1996) 37(1):29). Random cDNA clones from a tissue library are hybridized at moderate stringency to 300 7bp oligonucleotides. Each oligonucleotide has some measure of specific hybridization to that specific clone. The combination of 300 of these measures of hybridization for 300 probes equals the "hybridization signature" for a specific clone. Clones with similar sequence will have similar hybridization signatures. By developing a sorting/grouping algorithm to analyze these signatures, groups of clones in a library can be identified and brought together computationally. These groups of clones are termed "clusters". Depending on the stringency of the selection in the algorithm (similar to the stringency of hybridization in a

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classic library cDNA screening protocol), the "purity" of each cluster can be controlled. For example, artifacts of clustering may occur in computational clustering just as artifacts can occur in "wet-lab" screening of a cDNA library with 400 bp cDNA fragments, at even the highest stringency. The stringency used in the implementation of cluster herein provides groups of clones that are in general from the same cDNA or closely related cDNAs. Closely related clones can be a result of different length clones of the same cDNA, closely related clones from highly related gene families, or splice variants of the same cDNA.

Differential expression for a selected cluster was assessed by first determining the number of cDNA clones corresponding to the selected cluster in the first library (Clones in 1st), and the determining the number of cDNA clones corresponding to the selected cluster in the second library (Clones in 2nd). Differential expression of the selected cluster in the first library relative to the second library is expressed as a "ratio" of percent expression between the two libraries. In general, the "ratio" is calculated by: 1) calculating the percent expression of the selected cluster in the first library by dividing the number of clones corresponding to a selected cluster in the first library by the total number of clones analyzed from the first library; 2) calculating the percent expression of the selected cluster in the second library by dividing the number of clones corresponding to a selected cluster in a second library by the total number of clones analyzed from the second library; 3) dividing the calculated percent expression from the first library by the calculated percent expression from the second library. If the "number of clones" corresponding to a selected cluster in a library is zero, the value is set at 1 to aid in calculation. The formula used in calculating the ratio takes into account the "depth" of each of the libraries being compared, i.e., the total number of clones analyzed in each library.

In general, a polynucleotide is said to be significantly differentially expressed between two samples when the ratio value is greater than at least about 2, preferably greater than at least about 3, more preferably greater than at least about 5, where the ratio value is calculated using the method described above. The significance of differential expression is determined using a z score test (Zar, <u>Biostatistical Analysis</u>, Prentice Hall, Inc., USA, "Differences between Proportions," pp 296-298 (1974).

Example 5: Polynucleotides Differentially Expressed in High Metastatic Potential

Breast Cancer Cells Versus Low Metastatic Breast Cancer Cells

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A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high metastatic potential breast cancer tissue and low metastatic breast cancer cells. Expression of these sequences in breast cancer can be valuable in determining diagnostic, prognostic and/or treatment information. For example, sequences that are highly expressed in the high metastatic potential cells can be indicative of increased expression of genes or regulatory sequences involved in the metastatic process. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant more aggressive treatment. In another example, sequences that display higher expression in the low metastatic potential cells can be associated with genes or regulatory sequences that inhibit metastasis, and thus the expression of these polynucleotides in a sample may warrant a more positive prognosis than the gross pathology would suggest.

The differential expression of these polynucleotides can be used as a diagnostic marker, a prognostic marker, for risk assessment, patient treatment and the like. These polynucleofide sequences can also be used in combination with other known molecular and/or biochemical markers.

The following tables summarize polynucleotides that are differentially expressed between high metastatic potential breast cancer cells and low metastatic potential breast cancer cells.

Table 5.

Differentially expressed polynucleotides: Higher expression in high metastatic potential breast cancer (lib3) relative to low metastatic breast cancer cells (lib4)

SEQ ID NOS:	Sequence Name	Cluster ID	Lib3 clones	Lib4 clones	lib3/lib4	Zscore
NOS. 45	RTA00000197AR.f.12.1	3513	17	5	3.317240	2.287632
146	RTA00000185AF.a.19.2	5749	9	0	8.780930	2.629923
154	RTA00000196F.e.7.1	1039	10	2	4.878294	1.978215
159	RTA00000182AF.1.12.1	1027	41	17	2.353059	2.926571
165	RTA00000192AF.g.23.1	6455	6	0	5.853953	2.011224
174	RTA00000181AF.e.22.3	3442	17	4	4.146550	2.562391
183	RTA00000198AF.c.17.1	6923	6	0	5.853953	2.011224
364	RTA00000187AF.g.13.1	2991	10	1	9.756589	2.371428
366	RTA00000192AF.o.19.1	3549	10	1	9.756589	2.371428
387	RTA00000191AF.j.14.1	1002	42	20	2.048883	2.570309
496	RTA00000190AF.p.3.1	2378	34	0	33.17240	5.588184
510	RTA00000178AF.n.23.1	3298	12	1	11.70790	2.729313
512	RTA00000191AF.c.3.1	3549	10	1	9.756589	2.371428
529	RTA00000178AF.b.13.1	3114	9	1	8.780930	2.174815
560	RTA00000184AF.i.23.3	1577	25	3	8.130490	3.903813
606	RTA00000179AR.e.01.4	2493	33	9	3.577416	3.469507
		•	115			

115

SEQ ID NOS:	Sequence Name	Cluster ID	Lib3 clones	Lib4 clones	lib3/lib4	Zscore
644	RTA00000197F.i.12.1	3605	14	1	13.65922	3.050936
646	RTA000001971.1.12.1 RTA00000186AF.d.24.1	3114	9	1	8.780930	2.174815
754	RTA00000187AF.l.11.1	4482	14	3	4.553074	2.374769
875	RTA00000107H :::7111 RTA00000401F.m.02.1	1573	34	7	4.738914	3.982056
902	RTA00000422F.c.02.1	2902	18	5	3.512372	2.443314
921	RTA00000418F.m.19.1	8890	6	0	5.853953	2.011224
942	RTA00000351R.g.11.1	3077	17	4	4.146550	2.562391
1095	RTA00000408F.1.13.1	4423	12	i	11.70790	2.729313
1104	RTA00000404F.m.10.2	779	60	22	2.660887	3.974953
1131	RTA00000400F.k.22.1	2512	7	0	6.829612	2.235371
1170	RTA000004001.k.22.11 RTA00000340R.f.05.1	3202	18	3	5.853953	2.998867
1184	RTA00000340Kii.03.1 RTA00000422F.c.17.1	1360	26	11	2.306102	2.226876
1205	RTA000004221.6.17.1 RTA00000118A.a.23.1	3500	12	3	3.902635	2.018050
1354	RTA00000401F.k.14.1	211	121	43	2.745458	5.856098
2124	RTA00000191AF.j.14.1	1002	42	20	2.048883	2.570309
1535	RTA00000405F.l.11.1	2055	29	8	3.536763	3.213373
1751	RTA00000423F.j.03.1	5391	6	0	5.853953	2.011224
1764	RTA00000399F.o.24.1	2272	17	1	16.58620	3.483575
1777	RTA00000401F.j.15.1	3061	14	0	13.65922	3.428594
1795	RTA00000348R.o.12.1	2263	6	0	5.853953	2.011224
1869	RTA00000340F.f.22.1	1720	57	8	6.951569	5.855075
1882	RTA00000401F.g.22.1	1147	28	12	2.276537	2.294031
1890	RTA00000346F.o.16.1	176	170	44	3.769591	8.366611
1915	RTA00000400F.g.02.1	1508	21	5	4.097767	2.879196
2040	RTA00000527F.j.02.2	4896	11	0	10.73224	2.974502
2059	RTA00000528F.i.22.1	2478	17	5	3.317240	2.287632
2223	RTA00000528F.j.11.1	1070	26	6	4.227855	3.289393
2245	RTA00000527F.k.09.1	213	17	4	4.146550	2.562391
2300	RTA00000528F.b.03.1	2078	11	2	5.366124	2.174565
2325	RTA00000525F.d.13.1	349	77	1	75.12573	8.384408
2462	RTA00000528F.g.22.2	920	76	32		4.010278
2488	RTA00000528F.h.02.2	1701	18	4	4.390465	2.714073
2492	RTA00000528F.c.11.1	1701	18	4	4.390465	2.714073

Differentially expressed polynucleotides: Higher expression in low Table 6. metastatic breast cancer cells (lib4) relative to high metastatic potential breast cancer (lib3) Sequence Name Cluster ID Lib4 Lib 3 lib4/lib3 Zscore **SEQ ID** Clones NOS: Clones 4188 4 13 3.33108 1.99126 RTA00000177AR.n.8.1 15 1 8 8.19958 2.03713 40392 36 RTA00000181AF.p.4.3 0 11.2744 3.05623 RTA00000199F.f.08.2 12445 11 44 1.99126 3.33108 89 RTA00000177AF.n.8.3 4188 4 13 RTA00000186AF.p.09.2 6879 3 43 14.6909 5.83444 172 37 157 4.34910 8.71727 RTA00000201F.d.09.1 203 1827 RTA00000192AF.a.24.1 13183 0 7 7.17463 2.30057 261 2 20 10.2494 3.68254 RTA00000182AF.j.20.1 4769 419 2 20 10.2494 3.68254 420 RTA00000181AF.c.11.1 4769 2.45316 10.2494 503 RTA00000197AF.k.9.1 3138 1 10 552 RTA00000193AF.b.24.1 35 386 1967 5.22298 33.2328 1600 23 23.5738 4.64683 0 564 RTA00000200AF.g.18.1

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SEQ ID	Sequence Name	Cluster ID	Lib4	Lib 3	lib4/lib3	Zscore
NOS:	1		Clones	Clones		
570	RTA00000183AF.a.19.2	3788	0	6	6.14969	2.07158
590	RTA00000190AF.d.2.1	2444	26	55	2.16815	3.22244
693	RTA00000198F.m.12.1	4	987	2807	2.91492	30.3819
707	RTA00000179AF.p.15.1	5622	2	13	6.66216	2.62993
711	RTA00000198F.i.2.1	8076	0	9	9.22453	2.70385
726	RTA00000200R.f.10.1	4	987	2807	2.91492	30.3819
746	RTA00000178AF.i.01.2	4	987	2807	2.91492	30.3819
756	RTA00000404F.a.02.1	9738	1	13	13.3243	2.98623
990	RTA00000126A.o.23.1	6268	3	18	6.14969	3.11179
1122	RTA00000401F.o.06.1	2679	4	23	5.89345	3.52846
1142	RTA00000411F.a.15.1	73812	0	12	12.2993	3.21838
1286	RTA00000345F.n.12.1	7337	3	16	5.46639	2.80694
1289	RTA00000126A.g.7.1	1902	13	48	3.78442	4.45002
1435	RTA00000345F.e.11.1	4392	1	8	8.19958	2.03713
1860	RTA00000340F.p.18.1	287	6	173	29.5526	12.5749
1933	RTA00000400F.f.11.1	4088	0	82	84.0457	9.05778
1934	RTA00000341F.o.12.1	2883	9	21	2.39154	2.07600
1979	RTA00000122A.h.24.1	48	412	1020	2.53749	16.5262
1980	RTA00000346F.j.13.1	5337	5	17	3.48482	2.40321
2007	RTA00000400F.g.08.1	1275	15	32	2.18655	2.41857
2023	RTA00000523F.d.19.1	26489	1	8	8.19958	2.03713
2409	RTA00000526F.d.17.1	2757	4	16	4.09979	2.51500
1220	RTA00000528F.d.04.1	2395	12	37	3.16025	3.51521

Example 6: Polynucleotides Differentially Expressed in High Metastatic Potential Lung

Cancer Cells Versus Low Metastatic Lung Cancer Cells

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high metastatic potential lung cancer tissue and low metastatic lung cancer cells. Expression of these sequences in lung cancer tissue can be valuable in determining diagnostic, prognostic and/or treatment information. For example, sequences that are highly expressed in the high metastatic potential cells are associated can be indicative of increased expression of genes or regulatory sequences involved in the metastatic process. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant more aggressive treatment. In another example, sequences that display higher expression in the low metastatic potential cells can be associated with genes or regulatory sequences that inhibit metastasis, and thus the expression of these polynucleotides in a sample may warrant a more positive prognosis than the gross pathology would suggest.

The differential expression of these polynucleotides can be used as a diagnostic marker, a prognostic marker, for risk assessment, patient treatment and the like. These

polynucleotide sequences can also be used in combination with other known molecular and/or biochemical markers.

The following tables summarize polynucleotides that are differentially expressed between high metastatic potential lung cancer cells and low metastatic potential lung cancer cells:

Table 7 Differentially expressed polynucleotides: Higher expression in high metastatic potential lung cancer cells (lib8) relative to low metastatic lung cancer cells (lib9)

(1109)						
SEQ ID	Sequence Name	Cluster ID	Lib8	Lib9	lib8/lib9	Zscore
NO:	-		clones	clones		
10	RTA00000198AF.n.16.1	3721	9	0	12.5772	3.20845
54	RTA00000200F.o.22.1	983	8	1	11.1797	2.53243
65	RTA00000198AF.m.16.1	51	348	66	7.36849	17.4315
171	RTA00000198R.c.07.1	19181	6	0	8.38484	2.48169
203	RTA00000201F.d.09.1	1827	45	15	4.19242	5.09891
252	RTA00000181AF.e.18.3	8	1355	122	15.5211	39.0214
253	RTA00000181AF.e.17.3	8	1355	122	15.5211	39.0214
285	RTA00000181AR.j.14.3	5399	12	0	16.7696	3.80239
419	RTA00000182AF.j.20.1	4769	10	3	4.65824	2.29362
420	RTA00000181AF.c.11.1	4769	10	3	4.65824	2.29362
491	RTA00000196F.k.11.1	3	986	392	3.51507	22.4683
525	RTA00000198AF.c.7.1	19181	6	0	8.38484	2.48169
526	RTA00000185AF.e.20.1	5865	12	0	16.7696	3.80239
552	RTA00000193AF.b.24.1	35	868	11	110.273	34.2897
693	RTA00000198F.m.12.1	4	506	209	3.38335	15.7309
700	RTA00000183AF.i.18.2	40129	7	0	9.78231	2.74441
726	RTA00000200R.f.10.1	4	506	209	3.38335	15.7309
742	RTA00000177AF.m.1.1	14929	23	16	2.00886	2.02420
746	RTA00000178AF.i.01.2	4	506	209	3.38335	15.7309
861	RTA00000339F.f.11.1	5832	5	0	6.98736	2.18988
990	RTA00000126A.o.23.1	6268	5 '	0	6.98736	2.18988
1088	RTA00000399F.f.11.1	40167	8	0	11.1797	2.98512
1288	RTA00000423F.e.11.1	2566	11	2	7.68610	2.85611
1417	RTA00000339F.o.07.1	2566	11	2	7.68610	2.85611
1444	RTA00000419F.p.03.1	1937	10	3	4.65824	2.29362
1454	RTA00000340F.1.05.1	38935	7	0	9.78231	2.74441
1570	RTA00000403F.a.17.1	13686	8	0	11.1797	2.98512
1597	RTA00000401F.n.23.1	1552	8	1	11.1797	2.53243
1979	RTA00000122A.h.24.1	48	342	155	3.08345	12.2138
2024	RTA00000528F.b.23.1	1605	22	4	7.68610	4.23808
2034	RTA00000528F.m.16.1	4468	6	1	8.38484	1.97787
2126	RTA00000526F.d.01.1	4468	6	-1	8.38484	1.97787

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Table 8 Differentially expressed polynucleotides: Higher expression in low metastatic lung cancer cells (lib9) relative to high metastatic potntial lung cancer cells SEQ ID Sequence Name Cluster Lib8 Lib9 lib9/lib8 Zscore

SEQ ID	Sequence Name	Cluster	Lib8	Lib9	lib9/lib8	Zscore
NO:		ID	clones	clones		
174	RTA00000181AF.e.22.3	3442	5	23	3.291654	2.368262
254	RTA00000178AF.n.2.1	17083	0	8	5.724617	2.034117
466	RTA00000177AF.p.20.1	4141	4	27	4.830145	3.070829
571	RTA00000198AF.b.14.1	801	16	46	2.057284	2.411087
574	RTA00000192AF.f.3.1	5257	5	25	3.577885	2.596857
590	RTA00000190AF.d.2.1	2444	12	37	2.206362	2.299984
922	RTA00000399F.I.14.1	3354	5	20	2.862308	1.998763
1355	RTA00000406F.m.04.1	14959	11	41	2.667151	2.865855
1422	RTA00000405F.h.07.2	4984	3	16	3.816411	2.058861
2007	RTA00000400F.g.08.1	1275	10	42	3.005423	3.147111
2038	RTA00000527F.p.06.1	1292	8	33	2.951755	2.724411
2245	RTA00000527F.k.09.1	213	137	403	2.104945	7.661033

Example 7: Polynucleotides Differentially Expressed in High Metastatic Potential Colon Cancer Cells Versus Low Metastatic Colon Cancer Cells

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high metastatic potential colon cancer tissue and low metastatic colon cancer cells. Expression of these sequences in colon cancer tissue can be valuable in determining diagnostic, prognostic and/or treatment information. For example, sequences that are highly expressed in the high metastatic potential cells can be indicative of increased expression of genes or regulatory sequences involved in the metastatic process. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant more aggressive treatment. In another example, sequences that display higher expression in the low metastatic potential cells can be associated with genes or regulatory sequences that inhibit metastasis, and thus the expression of these polynucleotides in a sample may warrant a more positive prognosis than the gross pathology would suggest.

The differential expression of these polynucleotides can be used as a diagnostic marker, a prognostic marker, for risk assessment, patient treatment and the like. These polynucleotide sequences can also be used in combination with other known molecular and/or biochemical markers.

The following table summarizes identified polynucleotides with differential expression between high metastatic potential colon cancer cells and low metastatic potential colon cancer cells:

Table 9 Differentially expressed polynucleotides: Higher expression in high metastatic potential colon cancer (lib1) relative to low metastatic colon cancer cells (lib2)

metastatic potential colon cancer (1101) relative to low metastatic colon cancer cens (1102)								
SEQ ID NO:	Sequence Name	Cluster ID	Lib1 clones	Lib2 clones	lib1/lib2	Zscore		
228	RTA00000187AR.h.15.2	6660	7	0	6.489973399	2.169320547		
280	RTA00000193AF.b.18.1	7542	8	0	7.417112456	2.36964728		
355	RTA00000184AR.b.24.1	5777	9	1	8.344251513	2.09555146		
491	RTA00000196F.k.11.1	3	5268	2164	2.257009497	32.96556438		
603	RTA00000183AR.d.11.3	6420	8	0	7.417112456	2.36964728		
680	RTA00000177AF.f.10.1	6420	8	0	7.417112456	2.36964728		
752	RTA00000192AF.o.7.1	5275	11	2	5.099264814	2.083995588		
753	RTA00000192AF.o.17.1	5275	11	2	5.099264814	2.083995588		
1241	RTA00000346F.l.13.1	7542	8	0	7.417112456	2.36964728		
1264	RTA00000349R.g.10.1	5777	9	1	8.344251513	2.09555146		
1401	RTA00000421F.m.14.1	3524	21	6	3.2449867	2.499690198		
1442	RTA00000350R.g.10.1	9026	7	0	6.489973399	2.169320547		
1514	RTA00000399F.o.06.1	13574	7	0	6.489973399	2.169320547		
1851	RTA00000421F.a.06.1	2385	27	4	6.258188635	3.743586088		
1915	RTA00000400F.g.02.1	1508	46	17	2.508729213	3.230059264		
2024	RTA00000528F.b.23.1	1605	36	11	3.034273278	3.244010467		
2066	RTA00000528F.m.12.1	5768	12	0		3.046665462		

Table 10 Differentially expressed polynucleotides: Higher expression in low metastatic colon cancer cells (lib2)relative to high metastatic potential colon cancer (lib1) SEO ID Sequence Name Cluster Lib1 Lib2 lib2/lib1 Zecore

SEQ ID	Sequence Name	Cluster	Lib1	Lib2	lib2/lib1	Zscore
NOS:		ID	clones	clones		
33	RTA00000178AR.a.20.1	945	9	21	2.51670	2.21703
250	RTA00000192AF.j.21.1	2289	3	23	8.26916	3.92187
282	RTA00000193AF.c.15.1	3726	3	14	5.03340	2.58312
370	RTA00000179AF.c.15.3	2995	4	13	3.50540	2.09770
387	RTA00000191AF.j.14.1	1002	12	65	5.84234	6.26259
443	RTA00000197AR.i.17.1	3516	5	17	3.66719	2.52439
460	RTA00000179AF.c.15.1	2995	4	13	3.50540	2.09770
545	RTA00000196F.a.2.1	3575	5	14	3.02004	2.00158
560	RTA00000184AF.i.23.3	1577	12	40	3.59528	4.01991
703	RTA00000198F.l.09.1	3611	2	13	7.01081	2.73040
704	RTA00000190AF.o.12.1	3438	5	14	3.02004	2.00158
1095	RTA00000408F.1.13.1	4423	1	8	8.62869	2.11495
1104	RTA00000404F.m.10.2	779	27	54	2.15717	3.23169
1205	RTA00000118A.a.23.1	3500	3	13	4.67387	2.40298
1354	RTA00000401F.k.14.1	211	109	206	2.03843	6.08597
1387	RTA00000191AF.j.14.1	1002	12	65	5.84234	6.26259
1734	RTA00000345F.b.17.1	945	9	21	2.51670	2.21703
1742	RTA00000422F.b.22.1	2368	14	34	2.61942	3.00662
1954	RTA00000401F.j.23.1	570	59	148	2.70560	6.66631
2262	RTA00000527F.o.12.1	688	29	60	2.23155	3.53946
2325	RTA00000525F.d.13.1	349	69	138	2.15717	5.27497

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Example 8: Polynucleotides Differentially Expressed in High Metastatic Potential Colon Cancer Patient Tissue Versus Normal Patient Tissue

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high metastatic potential colon cancer tissue and normal tissue. Expression of these sequences in colon cancer tissue can be valuable in determining diagnostic, prognostic and/or treatment information. For example, sequences that are highly expressed in the high metastatic potential cells are associated can be indicative of increased expression of genes or regulatory sequences involved in the advanced disease state which involves processes such as angiogenesis, dedifferentiation, cell replication, and metastasis. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant more aggressive treatment.

The differential expression of these polynucleotides can be used as a diagnostic marker, a prognostic marker, for risk assessment, patient treatment and the like. These polynucleotide sequences can also be used in combination with other known molecular and/or biochemical markers.

The following tables summarize polynucleotides that are differentially expressed between high metastatic potential colon cancer cells and normal colon cells:

Table 11 Differentially expressed polynucleotides isolated from samples from two patients (UC#2 and UC#3): Higher expression in high metastatic potential colon tissue (UC#2:lib17; UC#3:lib20) vs. normal colon tissue (UC#2:lib15; UC#3:lib18)

SEQ ID NO:	Sequence Name	Cluster ID	lib15 clones	lib17 clones	lib17/lib15	Zscore
65	RTA00000198AF.m.16.1	51	1	10	9.27022	2.28830
1780	RTA00000118A.j.24.1	18	4	23	5.33037	3.27028
1899	RTA00000345F.j.09.1	13	14	80	5.29727	6.34580
SEQ ID NO:	Sequence Name	Cluster ID	lib18 clones	lib20 clones	lib20/lib18	Zscore
1899	RTA00000345F.j.09.1	13	12	23	2.24234	2.16077

Table 12 Differentially expressed polynucleotides isolated from samples from two patients (UC#2 and UC#3): Higher expression in normal colon tissue (UC#2:lib15; UC#3:lib18)vs. high metastatic potential colon tissue (UC#2:lib17; UC#3:lib20).

SEQ ID	Sequence Name	Cluster	Lib5	L1ib7	lib15/lib17	Z Score:
NO:		ID	Clones	Clones		>2.5899%; >1.96
491	RTA00000196F.k.11.1	3	242	26	10.04	13.78900072
SEO ID	Sequence Name	Cluster	Lib18	Lib20	lib18/lib20	Zscore

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NO: ID clones clones 491 RTA00000196F.k.11.1 3 409 46 7.59993 15.3998

Example 9: Polynucleotides Differentially Expressed in High Colon Tumor Potential Patient Tissue Versus Metastasized Colon Cancer Patient Tissue

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high tumor potential colon cancer tissue and cells derived from high metastatic potential colon cancer cells. Expression of these sequences in colon cancer tissue can be valuable in determining diagnostic, prognostic and/or treatment information associated with the transformation of precancerous tissue to malignant tissue. This information can be useful in the prevention of achieving the advanced malignant state in these tissues, and can be important in risk assessment for a patient.

The following table summarizes identified polynucleotides with differential expression between high tumor potential colon cancer tissue and cells derived from high metastatic potential colon cancer cells:

Table 13 Differentially expressed polynucleotides: High tumor potential colon tissue vs. metastatic colon tissue

SEQ ID	Sequence Name	Cluster ID	L19	L20	lib19/lib20	Zscore
NO:	-		clones	clones		
252	RTA00000181AF.e.18.3	8	14	1	10.4712	2.56699
253	RTA00000181AF.e.17.3	8	14	1	10.4712	2.56699
491	RTA00000196F.k.11.1	3	328	46	5.33318	11.8962
581	RTA00000191AF.p.3.2	17	24	2	8.97535	3.41950
693	RTA00000198F.m.12.1	4	26	8	2.43082	2.09705
726	RTA00000200R.f.10.1	4	26	8	2.43082	2.09705
746	RTA00000178AF.i.01.2	4	26	8	2.43082	2.09705
1780	RTA00000118A.j.24.1	18	80	13	4.60274	5.51440
1899	RTA00000345F.j.09.1	13	148	23	4.81287	7.68618

20 <u>Example 10</u>: <u>Polynucleotides Differentially Expressed in High Tumor Potential Colon</u> <u>Cancer Patient Tissue Versus Normal Patient Tissue</u>

A number of polynucleotide sequences have been identified that are differentially expressed between cells derived from high tumor potential colon cancer tissue and normal tissue. Expression of these sequences in colon cancer tissue can be valuable in determining diagnostic, prognostic and/or treatment information associated with the prevention of achieving the malignant state in these tissues, and can be important in risk assessment for a

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patient. For example, sequences that are highly expressed in the potential colon cancer cells are associated with or can be indicative of increased expression of genes or regulatory sequences involved in early tumor progression. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant closer attention or more frequent screening procedures to catch the malignant state as early as possible.

The following tables summarize polynucleotides that are differentially expressed between high metastatic potential colon cancer cells and normal colon cells:

Table 14 Differentially expressed polynucleotides detected in samples from two patients (UC#2 and UC#3): Higher expression in tumor potential colon tissue (UC#2:lib16; UC#3:lib19)vs. normal colon tissue (UC#2:lib15; UC#3:lib18)

SEQ ID	Sequence Name	Cluster	Lib15	Lib16	lib16/lib15	Zscore
NO:		ID	clones	clones		
1899	RTA00000345F.j.09.1	13	14	50	3.43709	4.22436
SEQ ID	Sequence Name	Cluster	Lib18	Lib19	lib19/lib18	Zscore
NO:		ID	clones	clones		
65	RTA00000198AF.m.16.1	51	0	14	12.2505	3.23250
252	RTA00000181AF.e.18.3	8	1	14	12.2505	2.84687
253	RTA00000181AF.e.17.3	8	1	14	12.2505	2.84687
581	RTA00000191AF.p.3.2	17	4	24	5.25021	3.24580
693	RTA00000198F.m.12.1	4	6	26	3.79182	2.98901
716	RTA00000200F.p.05.1	3984	0	7	6.12525	2.09621
726	RTA00000200R.f.10.1	4	6	26	3.79182	2.98901
746	RTA00000178AF.i.01.2	4	6	26	3.79182	2.98901
1780	RTA00000118A.j.24.1	18	10	80	7.00028	6.65963
1899	RTA00000345F.j.09.1	13	12	148	10.7921	9.86174

Table 15 Differentially expressed polynucleotides: Higher expression in normal colon tissue (UC#2:lib15) vs. tumor potential colon tissue (UC#2:lib16)

SEQ ID	Sequence Name	Cluster	Lib15	Lib16	lib15/lib16	Zscore
NO:		ID	clones	clones		
491	RTA00000196F.k.11.1	3	242	39	6.44765	12.3988

Example 11: Polynucleotides Differentially Expressed in Growth Factor-Stimulated Human Microvascular Endothelial Cells (HMEC) Relative to Untreated HMEC

A number of polynucleotide sequences have been identified that are differentially expressed between human microvascular endothelial cells (HMEC) that have been treated with growth factors relative to untreated HMEC.

Sequences that are differentially expressed between growth factor-treated HMEC and untreated HMEC can represent sequences encoding gene products involved in angiogenesis, metastasis (cell migration), and other development and oncogenic processes. For example, sequences that are more highly expressed in HMEC treated with growth factors (such as bFGF or VEGF) relative to untreated HMEC can serve as markers of

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cancer cells of higher metastatic potential. Detection of expression of these sequences in colon cancer tissue can be valuable in determining diagnostic, prognostic and/or treatment information associated with the prevention of achieving the malignant state in these tissues, and can be important in risk assessment for a patient. A patient sample displaying an increased level of one or more of these polynucleotides may thus warrant closer attention or more frequent screening procedures to catch the malignant state as early as possible.

The following table summarizes identified polynucleotides with differential expression between growth factor-treated and untreated HMEC.

10 **Table 16** Differentially expressed polynucleotides: Higher expression in bFGF treated HMEC (lib13) vs. untreated HMEC (lib12)

SEQ ID NO:	Sequence Name	 Lib12 clones	_	lib13/lib12	Zscore
648	RTA00000199F.i.9.1	 25	52	2.07199	2.94741

Table 17 Differentially expressed polynucleotides: Higher expression in VEGF treated HMEC (lib14) vs. untreated HMEC (lib12)

SEQ ID	Sequence Name	Cluster	Lib12	Lib14	lib14/lib12	Zscore
NO:		ID	clones	clones		
648	RTA00000199F.i.9.1	7	25	67	2.62449	4.17666
1899	RTA00000345F.j.09.1	13	22	49	2.18114	2.99887

Example 12: Polynucleotides Differentially Expressed Across Multiple Libraries

A number of polynucleotide sequences have been identified that are differentially expressed between cancerous cells and normal cells across all three tissue types tested (*i.e.*, breast, colon, and lung). Expression of these sequences in a tissue or any origin can be valuable in determining diagnostic, prognostic and/or treatment information associated with the prevention of achieving the malignant state in these tissues, and can be important in risk assessment for a patient. These polynucleotides can also serve as non-tissue specific markers of, for example, risk of metastasis of a tumor. The following table summarizes identified polynucleotides that were differentially expressed but without tissue typespecificity in the breast, colon, and lung libraries tested.

Table 18 Polynucleotides Differentially Expressed Across Multiple Library Comparisons

SEQ ID NO.	Cluster	Clones in 1st Lib	Clones in 2nd Lib	Ratio	Cell or Tissue Sample and Cancer State Compared
					(Z Score)

SEQ ID NO.	Cluster	Clones in 1st Lib	Clones in 2nd Lib	Ratio	Cell or Tissue Sample and Cancer State Compared (Z Score)
2024	1605	lib1	lib2	lib1/lib2	colon: high met > low met
		36	11	3.0342732	(3.2440104)
		lib8	lib9	lib8/lib9	lung: high met > low met
		22	4	7.6861036	(4.2380835)
65	51	lib8	lib9	lib8/lib9	lung: high met > low met
		348	66	7.3684960	(17.431560)
		lib18	lib19	lib19/lib18	pt #3 colon: tumor > normal
		0	14	12.250507	(3.2325073)
		lib15	lib17	lib17/lib15	pt #2 colon: met > normal
		1	10	9.2702249	(2.2883061)
174	3442	lib8	lib9	lib9/lib8	lung: low met > high met
		5	23	3.2916548	(2.3682625)
		lib3	lib4	lib3/lib4	breast: high met > low met
		17	4	4.1465504	(2.5623912)
203	1827	lib8	lib9	lib8/lib9	lung: high met > low met
		45	15	4.1924201	(5.0989192)
		lib3	lib4	lib4/lib3	breast: low met > high met
		37	157	4.3491051	(8.7172773)
2245	213	lib8	lib9	lib9/lib8	lung: low met > high met
		137	403	2.1049458	(7.6610331)
		lib3	lib4	lib3/lib4	breast: high met > low met
		17	4	4.1465504	(2.5623912)
990	6268	lib8	lib9	lib8/lib9	lung: high met > low met
		5	0	6.9873669	(2.1898837)
		lib3	lib4	lib4/lib3	breast: low met > high met
		3	18	6.1496901	(3.1117967)
252	8	lib8	lib9	lib8/lib9	lung: high met > low met
		1355	122	15.521118	(39.021411)
		lib19	lib20	lib19/lib20	pt. #3 colon: tumor > met
		14	1	10.471247	(2.5669948)
		lib18	lib19	lib19/lib18	pt #3 colon: tumor > normal
T		1	14	12.250507	(2.8468716)
253	8	lib8	lib9	lib8/lib9	lung: high met > low met
		1355	122	15.521118	(39.021411)
		lib19	lib20	lib19/lib20	pt. #3 colon: tumor > met
		14	1	10.471247	(2.5669948)
		lib18	lib19	lib19/lib18	pt #3 colon: tumor > normal
2225	240	1	14	12.250507	(2.8468716)
2325	349	lib3	lib4	lib3/lib4	breast: high met > low met
		77	1	75.125736	(8.3844087)
		lib1	lib2	lib2/lib1	colon: low met > high met
		69	138	2.1571737	(5.2749799)

SEQ ID NO.	Cluster	Clones in 1st Lib	Clones in 2nd Lib	Ratio	Cell or Tissue Sample and Cancer State Compared (Z Score)
1095	4423	lib3	lib4	lib3/lib4	breast: high met > low met
		12	1	11.707907	(2.7293134)
		lib1	lib2	lib2/lib1	colon: low met > high met
		1	8	8.6286948	(2.1149516)
1124	779	lib3	lib4	lib3/lib4	breast: high met > low met
		60	22	2,6608879	(3.9749537)
		lib1	lib2	lib2/lib1	colon: low met > high met
		27	54	2.1571737	(3.2316908)
387	1002	lib3	lib4	lib3/lib4	breast: high met > low met
		42	20	2.0488837	(2.5703094)
		lib1	lib2	lib2/lib1	colon: low met > high met
		12	65	5.8423454	(6.2625969)
419	4769	lib8	lib9	lib8/lib9	lung: high met > low met
		10	3	4.6582446	(2.2936274)
		lib3	lib4	lib4/lib3	breast: low met > high met
		2	20	10.249483	(3.6825426)
420	4769	lib8	lib9	lib8/lib9	lung: high met > low met
		10	3	4.6582446	(2.2936274)
		lib3	lib4	lib4/lib3	breast: low met > high met
		2	20	10.249483	(3.6825426)
1205	3500	lib3	lib4	lib3/lib4	breast: high met > low met
		12	3	3.9026356	(2.0180506)
		lib1	lib2	lib2/lib1	colon: low met > high met
		3	13	4.6738763	(2.4029818)
491	3	lib1	lib2	lib1/lib2	colon: high met > low met
		5268	2164	2.2570094	(32.965564)
		lib8	lib9	lib8/lib9	lung: high met > low met
		986	392	3.5150733	(22.468331)
		lib19	lib20	lib19/lib20	pt #3 colon: tumor > met
		328	46	5.3331820	(11.896271)
		lib18	lib20	lib18/lib20	pt #3 colon: normal > met
		409	46	7.5999342	(15.399861)
		lib15	lib17	lib15/lib17	pt#2 colon: normal > met
		242	26	10.04	(13.789000)
		lib15	lib16	lib15/lib16	pt#2 colon: normal > tumor
4		242	39	6.44765	12.39883
552	35	lib8	lib9	lib8/lib9	lung: high met > low met
		868	11	110.27335	(34.289704)
		lib3	lib4	lib4/lib3	breast: low met > high met
		386	1967	5.2229880	(33.232871)
560	1577	lib3	lib4	lib3/lib4	breast: high met > low met
		25	3	8.1304909	(3.9038139)

SEQ ID NO.	Cluster	Clones in 1st Lib	Clones in 2nd Lib	Ratio	Cell or Tissue Sample and Cancer State Compared (Z Score)
		lib1	lib2	lib2/lib1	colon: low met > high met
		12	40	3.5952895	(4.0199130)
581	17	lib19	lib20	lib19/lib20	pt #3 colon: tumor > met
		24	2	8.9753551	(3.4195074)
		lib18	lib19	lib19/lib18	pt #3 colon: tumor > normal
		4	24	5.2502174	(3.2458055)
590	2444	lib3	lib4	lib4/lib3	breast: low met > high met
		26	55	2.1681599	(3.2224421)
		lib8	lib9	lib9/lib8	lung: low met > high met
		12	37	2.2063628	(2.2999846)
1354	211	lib3	lib4	lib3/lib4	breast: high met > low met
		121	43	2.7454588	(5.8560985)
		lib1	lib2	lib2/lib1	colon: low met > high met
		109	206	2.0384302	(6.0859794)
1387	1002	lib3	lib4	lib3/lib4	breast: high met > low met
		42	20	2.0488837	(2.5703094)
		lib1	lib2	lib2/lib1	colon: low met > high met
		12	65	5.8423454	(6.2625969)
648	7	lib12	lib14	lib14/lib12	HMEC: VEGF > untreated
		25	67	2.6244913	(4.1766696)
		lib12	lib13	lib13/lib12	HMEC: bFGF > untreated
		25	52	2.0719962	(2.9474155)
693	4	lib8	lib9	lib8/lib9	lung: high met > low met
		506	209	3.3833566	(15.730912)
		lib3	lib4	lib4/lib3	breast: low met > high met
		987	2807	2.9149240	(30.381945)
		lib19	lib20	lib19/lib20	pt#3 colon: tumor > met
		26	8	2.4308253	(2.0970580)
		lib18	lib19	lib19/lib18	pt#3 colon: tumor > normal
		6	26	3.7918237	(2.9890107)
726	4	lib8	lib9	lib8/lib9	lung: high met > low met
		506	209	3.3833566	(15.730912)
		lib3	lib4	lib4/lib3	breast: low met > high met
		987	2807	2.9149240	(30.381945)
		lib19	lib20	lib19/lib20	pt#3 colon: tumor > met
		26	8	2.4308253	(2.0970580)
		lib18	lib19	lib19/lib18	pt#3 colon: tumor > normal
		6	26	3.7918237	(2.9890107)
746	4	lib8	lib9	lib8/lib9	lung: high met > low met
		506	209	3.3833566	(15.730912)
		lib3	lib4	lib4/lib3	breast: low met > high met
		987	2807	2.9149240	(30.381945)

SEQ ID NO.	Cluster	Clones in 1st Lib	Clones in 2nd Lib	Ratio	Cell or Tissue Sample and Cancer State Compared (Z Score)
•		lib19	lib20	lib19/lib20	pt#3 colon: tumor > met
		26	8	2.4308253	(2.0970580)
		lib18	lib19	lib19/lib18	pt#3 colon: tumor > normal
		6	26	3.7918237	(2.9890107)
1780	18	lib19	lib20	lib19/lib20	pt#3 colon: tumor > met
		80	13	4.6027462	(5.5144093)
		lib18	lib19	lib19/lib18	pt#3 colon: tumor > normal
		10	80	7.0002899	(6.6596394)
		lib15	lib17	lib17/lib15	pt#3 colon: met > normal
		4	23	5.3303793	(3.2702852)
1899	13	lib19	lib20	lib19/lib20	pt#3 colon: tumor > met
		148	23	4.8128716	(7.6861840)
		lib18	lib20	lib20/lib18	pt#3 colon: met > normal
		12	23	2.2423439	(2.1607719)
		lib18	lib19	lib19/lib18	pt#3 colon: tumor > normal
		12	148	10.792113	(9.8617485)
		lib15	lib17	lib17/lib15	pt#2 colon: met > normal
		14	80	5.2972714	(6.3458044)
		lib15	lib16	lib16/lib15	pt#2 colon: tumor > normal
		14	50	3.4370927	(4.2243697)
		lib12	lib14	lib14/lib12	HMEC: VEGF > untreated
		22	49	2.1811410	(2.9988774)
1915	1508	lib1	lib2	lib1/lib2	colon: high met > low met
		46	17	2.5087292	(3.2300592)
		lib3	lib4	lib3/lib4	breast: high met > low met
		21	5	4.0977674	(2.8791960)
1979	48	lib8	lib9	lib8/lib9	lung: high met > low met
		342	155	3.0834574	(12.213852)
		lib3	lib4	lib4/lib3	breast: low met > high met
		412	1020	2.5374934	(16.526285)
2007	1275	lib3	lib4	lib4/lib3	breast: low met > high met
		15	32	2.1865564	(2.4185764)
		lib8	lib9	lib9/lib8	lung: low met > high met
		10	42	3.0054239	3.1471113

high met = high metastatic potential; low met = low metastatic potential; met = metastasized; tumor = non-metastasized tumor; pt = patient; #2 = UC#2; #3 = UC#3;

HMEC = human microvascular endothelial cell;

5 bFGF = bFGF treated; VEGF = VEGF treated

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Example 12: Polynucleotides Exhibiting Colon-Specific Expression

The cDNA libraries described herein were also analyzed to identify those polynucleotides that were specifically expressed in colon cells or tissue, *i.e.*, the polynucleotides were identified in libraries prepared from colon cell lines or tissue, but not in libraries of breast or lung origin. The polynucleotides that were expressed in a colon cell line and/or in colon tissue, but were present in the breast or lung cDNA libraries described herein, are shown in Table 19 (inserted before claims).

No clones corresponding to the colon-specific polynucleotides in the table above were present in any of Libraries 3, 4, 8, 9, 12, 13, 14, or 15. The polynucleotide provided above can be used as markers of cells of colon origin, and find particular use in reference arrays, as described above.

Example 13: Identification of Contiguous Sequences Having a Polynucleotide of the Invention

The novel polynucleotides were used to screen publicly available and proprietary databases to determine if any of the polynucleotides of SEQ ID NOS:1-2502 would facilitate identification of a contiguous sequence, *e.g.*, the polynucleotides would provide sequence that would result in 5' extension of another DNA sequence, resulting in production of a longer contiguous sequence composed of the provided polynucleotide and the other DNA sequence(s). Contiging was performed using the Gelmerge application (default settings) of GCG from the Univ. of Wisconsin.

Using these parameters, 146 contiged sequences were generated. These contiged sequences are provided as SEQ ID NOS:5107-5252 (see Table 1). The contiged sequences can be correlated with the sequences of SEQ ID NOS:1-2502 upon which the contiged sequences are based by, for example, identifying those sequences of SEQ ID NOS:1-2502 and the contiged sequences of SEQ ID NOS:5107-5252 that share the same clone name in Table 1.

The contiged sequences (SEQ ID NO:5107-5252) thus represent longer sequences that encompass a polynucleotide sequence of the invention. The contiged sequences were then translated in all three reading frames to determine the best alignment with individual sequences using the BLAST programs as described above for SEQ ID NOS:1-2502 and the validation sequences "SEQ ID NOS:2503-5106." Again the sequences were masked using the XBLAST program for masking low complexity as described above in Example 1

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(Table 2). Several of the contiged sequences were found to encode polypeptides having characteristics of a polypeptide belonging to a known protein families (and thus represent new members of these protein families) and/or comprising a known functional domain (Table 20). Thus the invention encompasses fragments, fusions, and variants of such polynucleotides that retain biological activity associated with the protein family and/or functional domain identified herein.

 Table 20
 Profile hits using contiged sequences

SEQ ID NO	Biological Activity (Profile)	Start	Stop	Score	Direction	Sequence Name
5111	7tm_2	71	915	8090	for	RTA00000399F.o.01.1
5120	7tm_2	101	919	8475	rev	RTA00000341F.m.21.1
5174	7tm_2	3	963	9431	for	RTA00000192AF.h.19.1
5197	7tm_2	214	1073	8528	rev	RTA00000192AF.f.3.1
5208	ANK	546	629	4920	for	RTA00000190AF.f.5.1
5120	asp	126	1067	6620	rev	RTA00000341F.m.21.1.
5241	asp	112	1094	6553	for	RTA00000418F.i.06.1
5243	asp	347	1028	5981	for	RTA00000339F.b.02.1
5197	ATPases	113	781	5690	for	RTA00000192AF.f.3.1
5239	ATPases	1	348	15955	for	RTA00000401F.m.07.1
5241	ATPases	110	823	6782	for	RTA00000418F.i.06.1
5243	ATPases	338	874	5832	for	RTA00000339F.b.02.1
5125	protkinase	59	685	5791	for	RTA00000182AF.c.5.1
5217	protkinase	75	1035	5405	for	RTA00000181AF.p.12.3
5237	protkinase	25	546	5107	rev	RTA00000118A.n.5.1
5248	protkinase	14	422	5103	rev	RTA00000419F.k.05.1
5252	protkinase	89	755	5499	for	RTA00000404F.m.17.2
5120	Wnt_dev_sign	3	948	11036	for	RTA00000341F.m.21.1
					_	

All stop/start sequences are provided in the forward direction.

Descriptions of the profiles for the indicated protein families and functional domains are provided in Example 3 above.

Those skilled in the art will recognize, or be able to ascertain, using not more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. Such specific embodiments and equivalents are intended to be encompassed by the following claims.

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All publications and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference. The citation of any publication is for its disclosure prior to the filing date and should not be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it is readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

Deposit Information:

The following materials were deposited with the American Type Culture

15 Collection: CMCC = (Chiron Master Culture Collection)

Cell Lines Deposited with ATCC

Deposit Date	ATCC Accession No.	CMCC Accession No.
March 19, 1998	CRL-12496	11606
May 15, 1998	CRL-12533	11611
May 15, 1998	CRL-12532	10583
October 9, 1998	CRL-12584	10377
	March 19, 1998 May 15, 1998 May 15, 1998	March 19, 1998 CRL-12496 May 15, 1998 CRL-12533 May 15, 1998 CRL-12532

cDNA Libraries Deposited with ATCC

cDNA Library No.	cDNA Library ES21	cDNA Library ES22	cDNA Library ES23
Deposit Date	January 22, 1999	January 22, 1999	January 22, 1999
ATCC Accession No.	ATCC No.	ATCC No.	ATCC No.
Clone Names	M00001575D:G05	M00001364A:E11	M00001489B:A06
	M00001460A:A03	M00001694C:H10	M00001585A:D06
	M00001655C:E04	M00003841D:E03	M00001637B:E07
	M00001676C:C11	M00004176D:B12	M00001529D:H02
	M00001679D:D05	M00001387B:E02	M00001500C:C08
	M00001546B:C05	M00004282B:A04	M00001483B:D03
	M00001453B:E10	M00001376B:F03	M00001623C:H07
		M00001445D:A06	M00003975B:F03
		M00001399C:H12	
		M00004208D:H08	

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cDNA Library No.	cDNA Library ES24	cDNA Library ES25	cDNA Library ES26
Deposit Date	January 22, 1999	January 22, 1999	January 22, 1999
ATCC Accession No.	ATCC No.	ATCC No.	ATCC No.
Clone Names	M00003987D:D06	M00001675D:B08	M00001479C:F10
	M00004073A:H12	M00001589B:E12	M00003842D:F08
	M00004104B:F11	M00001607D:A11	M00003901A:C09
	M00004237D:D08	M00001636A:E07	M00003982A:B06
	M00004111D:B07	M00001530A:B12	M00003824A:A06
	M00004138B:B11	M00001495B:B08	M00003845D:C03
	M00001391C:C04	M00001487C:F01	M00003856A:B07
	M00001448D:E12	M00001644B:D06	M00004104B:A02
	M00001450A:B03	M00003751C:A04	M00004110C:E03
	M00001451B:F01		

In addition, libraries of selected clones were deposited. The details of these deposits are provided in Tables 21-24.

This deposit is provided merely as convenience to those of skill in the art, and is not an admission that a deposit is required under 35 U.S.C. §112. The sequence of the polynucleotides contained within the deposited material, as well as the amino acid sequence of the polypeptides encoded thereby, are incorporated herein by reference and are controlling in the event of any conflict with the written description of sequences herein. A license may be required to make, use, or sell the deposited material, and no such license is granted hereby.

Retrieval of Individual Clones from Deposit of Pooled Clones

Where the ATCC deposit is composed of a pool of cDNA clones, the deposit was prepared by first transfecting each of the clones into separate bacterial cells. The clones were then deposited as a pool of equal mixtures in the composite deposit. Particular clones can be obtained from the composite deposit using methods well known in the art. For example, a bacterial cell containing a particular clone can be identified by isolating single colonies, and identifying colonies containing the specific clone through standard colony hybridization techniques, using an oligonucleotide probe or probes designed to specifically hybridize to a sequence of the clone insert (*e.g.*, a probe based upon unmasked sequence of the encoded polynucleotide having the indicated SEQ ID NO). The probe should be designed to have a T_m of approximately 80°C (assuming 2°C for each A or T and 4°C for each G or C). Positive colonies can then be picked, grown in culture, and the recombinant clone isolated. Alternatively, probes designed in this manner can be used to PCR to isolate a nucleic acid molecule from the pooled clones according to methods well known in the art,

e.g., by purifying the cDNA from the deposited culture pool, and using the probes in PCR reactions to produce an amplified product having the corresponding desired polynucleotide sequence.

Table 1.					
SEQ ID	Filing	SEQ ID	Sequence Name	Clone Name	Cluster
NO:	Date of	NO: in	1		ID
	Priority	Priority			
	Appln	Appln			
1	1/28/98	1	RTA00000197AF.i.16.1	M00001490A:D11	16402
2	1/28/98	2	RTA00000188AF.n.15.1	M00003804A:H04	0
3	1/28/98	3	RTA00000197AF.e.24.1	M00001456B:F10	39250
4	1/28/98	4	RTA00000198R.f.04.1	M00001607D:F07	5023
5	1/28/98	5	RTA00000195R.c.11.1	M00003811A:E03	66087
6	1/28/98	6	RTA00000195AF.c.16.1	M00003829C:A11	23508
7	1/28/98	7	RTA00000197AR.e.12.1	M00001454B:G07	22095
8	1/28/98	8	RTA00000200AF.h.11.2	M00004146A:C08	8399
9	1/28/98	9	RTA00000177AF.g.22.1	M00001347C:G08	7031
10	1/28/98	10	RTA00000198AF.n.16.1	M00001694C:H10	3721
11	1/28/98	11	RTA00000199AF.i.17.1	M00003880C:F10	9615
12	1/28/98	12	RTA00000183AF.i.15.2	M00001529B:C04	2642
13	1/28/98	13	RTA00000190AF.i.5.1	M00003919A:A10	0
14	1/28/98	14	RTA00000196R.c.11.2	M0000331371:110	13658
15	1/28/98	15	RTA00000177AR.n.8.1	M00001356D:F06	4188
16	1/28/98	16	RTA000001777Killio11	M00001350D:100	39252
17	1/28/98	17	RTA0000183AR.e.14.2	M00001506B:D09	17437
18	1/28/98	18	RTA00000195/AR.c.14.2	M00001350B:B09	39602
19	1/28/98	19	RTA00000185AF.a.8.1	M00001532C:100	4868
20	1/28/98	20	RTA00000181AF.l.14.2	M00001370D://05	2364
21	1/28/98	21	RTA00000131A.g.19.2	M000014349A:G10	36535
22	1/28/98	22	RTA00000137A.g.13.2	M00001718D:F07	8984
23	1/28/98	23	RTA000001977R.b.1	M00001710D:107	22636
23	1/28/98	24	RTA00000130R.5.00.1	M00004111D:D11	0
25	1/28/98	25	RTA00000196AF.c.1.1	M00001349C:C05	8171
26	1/28/98	26	RTA00000200R.g.09.1	M00004131B:H09	22785
27	1/28/98	27	RTA00000192AF.i.12.1	M00004169C:C12	5319
28	1/28/98	28	RTA00000178AR.o.01.5	M00001387B:H07	0
29	1/28/98	29	RTA00000200AF.b.19.1	M00004042D:H02	22847
30	1/28/98	30	RTA00000184AR.n.07.2	M00001561C:F06	0
31	1/28/98	31	RTA00000200F.m.15.1	M00004236C:D10	22601
32	1/28/98	32	RTA00000198R.m.19.1	M00001680D:D02	40041
33	1/28/98	33	RTA00000178AR.a.20.1	M00001362C:H11	945
34	1/28/98	34	RTA00000197AF.n.8.1	M00001536D:A12	4101
35	1/28/98	35	RTA00000191AF.n.17.1	M00004091B:D11	7848
36	1/28/98	36	RTA00000181AF.p.4.3	M00001460A:A03	40392
37	1/28/98	37	RTA00000181AF.n.15.2	M00001457A:B07	86128
38	1/28/98	38	RTA00000196R.k.07.1	M00001399C:D09	22443
39	1/28/98	39	RTA00000189AR.b.19.1	M00003832B:E01	5294
40	1/28/98	40	RTA00000200AR.e.02.1	M00004090A:F09	36059
41	1/28/98	41	RTA00000184F.k.12.1	M00001557D:D09	8761
42	1/28/98	42	RTA00000184F.j.21.1	M00001557A:D02	7065
43	1/28/98	43	RTA00000179AF.c.14.3	M00001392D:H04	0
44	1/28/98	44	RTA00000199F.f.08.2	M00003841D:E03	12445
45	1/28/98	45	RTA00000197AR.f.12.1	M00001458C:E01	3513
46	1/28/98	46	RTA00000182AF.f.13.1	M00001470C:B10	8010
47	1/28/98	47	RTA00000192AF.m.12.1	M00004191D:B11	0
48	1/28/98	48	RTA00000177AR.a.23.5	M00001339D:G02	6995
49	1/28/98	49	RTA00000198R.o.05.1	M00003750A:D01	26702

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority Appln	Priority Appln			
50	1/28/98	50	RTA00000201R.a.02.1	M00004295B:D02	35362
51	1/28/98	51	RTA00000199R.k.07.1	M00004273B:D02	12973
52	1/28/98	52	RTA00000201R.b.02.1	M00003301C:/103	22660
53	1/28/98	53	RTA00000199AF.p.9.1	M00004919B:G09	10430
54	1/28/98	54	RTA00000200F.o.22.1	M0000330071:210	983
55	1/28/98	55	RTA00000186AF.i.21.1	M00001232D:H01	6033
56	1/28/98	56	RTA00000177AF.e.9.1	M00001343D:C04	37442
57	1/28/98	57	RTA00000198AF.k.20.1	M00001660C:B12	22553
58	1/28/98	58	RTA00000199F.b.01.2	M00003778A:D08	19118
59	1/28/98	59	RTA00000195AF.b.13.1	M00001560D:A03	12605
59	2/24/98	78	RTA00000195AF.b.13.1	M00001560D:A03	12605
60	1/28/98	60	RTA00000196AR.i.12.3	M00001389D:G11	38800
61	1/28/98	61	RTA00000197AF.h.11.1	M00001476D:G03	22264
62	1/28/98	62	RTA00000190AF.a.18.2	M00003900D:B10	0
63	1/28/98	63	RTA00000184AF.k.19.1	M00001558B:D08	8022
64	1/28/98	64	RTA00000198AF.p.12.1	M00003763D:E10	8878
65	1/28/98	65	RTA00000198AF.m.16.1	M00001679D:D05	51
66	1/28/98	66	RTA00000199F.c.09.2	M00003800A:C09	16824
67	1/28/98	67	RTA00000200AF.g.07.1	M00004128B:G01	0
68	1/28/98	68	RTA00000184F.k.19.1	M00001558B:D08	8022
69	1/28/98	69	RTA00000186AF.h.8.1	M00001632C:C09	35547
70	1/28/98	70	RTA00000192AF.e.3.1	M00004138B:H02	13272
71	1/28/98	71	RTA00000193AR.o.16.3	M00004409B:A11	52972
72	1/28/98	72	RTA00000200F.a.6.1	M00004029B:F11	36952
73	1/28/98	73	RTA00000177AF.e.21.3	M00001344A:H07	4306
74	1/28/98	74	RTA00000196AF.h.20.1	M00001385B:F10	0
75	1/28/98	75	RTA00000180AR.h.19.2	M00001428A:H10	84182
76	1/28/98	76	RTA00000200AF.h.05.2	M00004142D:E10	10950
77	1/28/98	77	RTA00000197AF.n.2.1	M00001535A:D02	6229
78	1/28/98	78	RTA00000199R.f.09.1	M00003842B:D09	22907
79	1/28/98	79	RTA00000199AF.p.4.1	M00003985C:F01	10282
80	1/28/98	80	RTA00000196AF.p.13.2	M00001432A:E06	6125
81	1/28/98	81	RTA00000196AF.b.15.1	M00001347B:E01	5102
82	1/28/98	82	RTA00000183AF.1.18.1	M00001535D:C01	3484
83	1/28/98	83	RTA00000186AF.f.24.2	M00001629B:E06	0
84	1/28/98	84	RTA00000191AF.h.14.1	M00004056B:D09	13553
85 86	1/28/98	8 5	RTA00000200R.o.03.1	M00004257C:H06	22807
86	1/28/98	86	RTA00000189AF.I.22.1	M00003879C:G10	33333
87 87	2/24/98 1/28/98	245 8 7	RTA00000195AF.d.20.1	M00004117A:D11	37574
87 88	1/28/98	88	RTA00000195AF.d.20.1 RTA00000197AF.e.23.1	M00004117A:D11 M00001456B:C09	37574
89	1/28/98	89	RTA00000197AF.e.23.1 RTA00000177AF.n.8.3	M00001456D:C09	37157 41 88
90	1/28/98	90	RTA00000177AF.II.8.3 RTA00000199F.f.15.2	M00001336D:F06 M00003845A:H12	8772
91	1/28/98	91	RTA000001991.1.13.2 RTA00000198AF.j.19.1	M00003843A:1112 M00001653C:F12	38914
92	1/28/98	92	RTA00000198AF.j.18.1	M00001653B:G07	22759
93	1/28/98	93	RTA00000198A1.J.18.1	M00001033B:G07 M00004270A:F11	0
94	1/28/98	94	RTA000002001.0.11.1 RTA00000195AF.b.4.1	M00004270A:F11 M00001490C:D07	0
95	1/28/98	95	RTA00000175A1.b.4.1 RTA00000180AF.g.3.1	M00001490C:D07	9024
96	1/28/98	96	RTA00000197AF.j.20.1	M0000142571:C11	4915
97	1/28/98	97	RTA00000197AF.o.2.1	M00001470C:E11	5739
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SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
98	Appln 1/28/98	Appln 98	RTA00000200AF.f.14.1	M00004115D:C08	22051
90 99	1/28/98	99	RTA00000200AF.I.14.1 RTA00000184AF.d.8.1	M00004113D:C00	4393
100	1/28/98	100	RTA00000104AI .d.6.1 RTA00000200R.f.14.1	M0000134674:7000 M00004115D:C08	22051
100	1/28/98	100	RTA00000200R.1.14.1 RTA00000191AF.d.08.2	M00004113B:C00 M00003997B:G07	970
101	1/28/98	101	RTA00000191A1.d.00.2	M000033371D:G07	37844
102	1/28/98	102	RTA00000199K.j.00.1 RTA00000199F.e.10.1	M00003804D:G07	22906
103	1/28/98	103	RTA000001991.c.10.1	M0000302271102 M00001381A:D02	6636
104	1/28/98	105	RTA00000179AF.g.12.3	M00001391A:G03	36390
105	1/28/98	106	RTA00000177AF.n.21.1	M00001540B:C09	0
100	1/28/98	107	RTA00000197712111	M00001310B:209	9857
107	1/28/98	107	RTA00000130R.h.23.2	M00001528A:F09	18957
108	1/28/98	100	RTA00000197AF.d.12.1	M0000132071109 M00001451D:C10	39546
110	1/28/98	110	RTA00000197R1.d.12.1	M00001431D:C10	13075
111	1/28/98	111	RTA00000197R.ii.01.11	M000014707X:H01	22038
112	1/28/98	112	RTA00000176A1.0.12.1	M00003751D:D02	8010
113	1/28/98	113	RTA00000177AF.iii.8.1	M00001354E:E10	16934
113	1/28/98	113	RTA00000190A1.d.03.11 RTA00000200R.f.02.1	M00001334B:B10	7138
115	1/28/98	115	RTA00000200R.1.02.1 RTA00000179AR.o.20.3	M00004100A:A07	2409
116	1/28/98	116	RTA00000173AR.0.20.3 RTA00000181AR.k.24.3	M00001454B:C12	7005
117	1/28/98	110	RTA00000181AR.k.24.5 RTA00000199AF.j.18.1	M00001434D:C12 M00003889D:B09	5140
117	1/28/98	117	RTA00000199A1.J.18.1 RTA00000199F.b.24.2	M00003794A:B03	0
118	1/28/98	119	RTA000001991.0.24.2 RTA00000181AR.k.24.2	M00003794A:B03 M00001454B:C12	7005
119	1/28/98	120	RTA00000181AR.k.24.2 RTA00000178AR.m.19.5	M00001434D:C12	0
121	1/28/98	120	RTA00000178AK.iii.19.5	M00001304D:1107 M00003979A:F03	16721
121	1/28/98	121	RTA00000197AF.I.15.1	M00003575A:103	4947
123	1/28/98	123	RTA00000197AF.k.6.1	M00001917B:G00 M00004078B:A11	5451
123	1/28/98	123	RTA00000191AI.R.0.1	M00003933C:D06	19122
125	1/28/98	125	RTA00000197AF.k.15.1	M00003535C:D00	22750
126	1/28/98	126	RTA00000201F.d.16.1	M00004388B:A08	0
127	1/28/98	127	RTA00000178AF.k.18.1	M00001382A:F04	9755
128	1/28/98	128	RTA00000196F.i.12.1	M00001389D:G11	38800
129	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
130	1/28/98	130	RTA00000196AF.h.23.1	M00001386A:C02	13357
131	1/28/98	131	RTA00000185AF.d.11.2	M00001579D:C03	6539
132	1/28/98	132	RTA00000178AF.f.20.3	M00001372C:F07	39881
133	1/28/98	133	RTA00000181AR.n.20.3	M00001457B:E03	0
134	1/28/98	134	RTA00000197F.e.11.1	M00001454B:G03	2306
135	1/28/98	135	RTA00000196AF.c.22.1	M00001352D:C05	22548
136	1/28/98	136	RTA00000197AF.c.10.1	M00001448B:F06	10400
137	1/28/98	137	RTA00000181AF.m.4.3	M00001455A:E09	13238
138	1/28/98	138	RTA00000182AF.a.3.3	M00001462B:A10	0
139	1/28/98	139	RTA00000191AF.d.01.2	M00003996A:A06	7031
140	1/28/98	140	RTA00000199F.a.2.1	M00003772A:D07	12674
141	1/28/98	141	RTA00000196AF.c.6.1	M00001350A:D06	23148
142	1/28/98	142	RTA00000198AF.k.19.1	M00001660B:C04	75879
143	1/28/98	143	RTA00000199R.h.09.1	M00003867C:H09	76020
144	1/28/98	144	RTA00000198AF.o.18.1	M00003755A:A09	13018
145	1/28/98	145	RTA00000178AF.h.24.1	M00001376B:C06	6745
146	1/28/98	146	RTA00000185AF.a.19.2	M00001571C:H06	5749
147	1/28/98	147	RTA00000185AF.c.24.2	M00001578B:E04	23001

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
1.40	Appln	Appln	DT40000100F L 17.2	MANNON 2071 A . A 05	36254
148	1/28/98	148	RTA00000199F.h.17.2	M00003871A:A05 M00001450D:D04	30234 87226
149	1/28/98	149 150	RTA00000181AR.h.06.3	M00001450D.D04 M00001557C:H07	7065
150	1/28/98	150	RTA00000184F.k.09.1 RTA00000200R.l.17.1	M00001337C:H07 M00004217C:D03	12771
151	1/28/98	151		M00004217C.D03 M00001352C:H02	8934
152	1/28/98	152	RTA00000196AF.c.20.1 RTA00000200F.n.17.2	M00001332C:H02 M00004252C:E03	19064
153	1/28/98	153	RTA00000200F.fl.17.2 RTA00000196F.e.7.1	M00004232C.E03 M00001360D:E11	1039
154 155	1/28/98 1/28/98	154	RTA00000196F.e.7.1 RTA00000197F.e.8.1	M00001300D:E11	3135
156	1/28/98	156	RTA00000197F.e.8.1 RTA00000199R.o.12.1	M00001434A.C11 M00003977A:E04	16128
		157	RTA00000199R.0.12.1 RTA00000188AF.n.01.1	M00003977A:E04 M00003801A:B10	36412
157 158	1/28/98 1/28/98	157	RTA00000198AF.k.03.1	M00003801A.B10 M00001655A:F06	22765
158	1/28/98	159	RTA00000198AF.k.03.1 RTA00000182AF.l.12.1	M00001033A:100	1027
160	1/28/98	160	RTA00000192AF.h.12.1 RTA00000192AF.b.20.1	M00001487A.A03 M00004118D:E08	0
161	1/28/98	161	RTA00000192AF.b.20.1 RTA00000183AF.e.23.2	M00004118D:L08	0
162	1/28/98	162	RTA00000163AF.e.23.2 RTA00000201F.e.15.1	M00001300D:A09	9960
163	1/28/98	163	RTA000002011.e.13.1 RTA00000192AR.e.13.3	M00004444B:B11	9457
164	1/28/98	164	RTA00000192AR.e.13.5 RTA00000193AR.i.14.4	M00004142A:B12	9457
165	1/28/98	165	RTA00000193AR.i.14.4 RTA00000192AF.g.23.1	M00004307C:A00	6455
166	1/28/98	166	RTA00000192AF.g.23.1 RTA00000198AF.f.21.1	M00004137C:A09	22676
167	1/28/98	167	RTA00000179AF.d.22.3	M00001394C:C11	7955
168	1/28/98	168	RTA00000177AR.k.23.1	M00001354C:C11	35550
169	1/28/98	169	RTA000001777R.R.25.1	M00001332D:D02	8685
170	1/28/98	170	RTA00000197AF.d.23.1	M00001360C:102	16130
171	1/28/98	171	RTA00000198R.c.07.1	M00001575D:G05	19181
172	1/28/98	172	RTA00000186AF.p.09.2	M00001655C:E04	6879
173	1/28/98	173	RTA00000200AR.b.07.1	M00004039C:C01	17125
174	1/28/98	174	RTA00000181AF.e.22.3	M00001448D:F09	3442
175	1/28/98	175	RTA00000200F.i.5.1	M00004156B:A12	22892
176	1/28/98	176	RTA00000183AF.h.19.1	M00001528A:A01	5175
177	1/28/98	177	RTA00000197AF.c.3.1	M00001447C:C01	3145
178	1/28/98	178	RTA00000200F.o.03.1	M00004257C:H06	22807
179	1/28/98	179	RTA00000179AF.f.20.3	M00001397B:B09	16154
180	1/28/98	180	RTA00000199AF.j.12.1	M00003887A:A06	22461
181	1/28/98	181	RTA00000198AF.d.2.1	M00001585A:F07	0
182	1/28/98	182	RTA00000196AF.h.16.1	M00001384C:E03	39895
183	1/28/98	183	RTA00000198AF.c.17.1	M00001579C:E08	6923
184	1/28/98	184	RTA00000197AF.f.7.1	M00001457C:C11	19261
185	2/24/98	234	RTA00000195AF.d.4.1	M00003881D:D06	22766
185	1/28/98	185	RTA00000195AF.d.4.1	M00003881D:D06	22766
186	1/28/98	186	RTA00000198R.p.09.1	M00003761D:E02	10473
187	1/28/98	187	RTA00000180AR.j.04.4	M00001429C:G12	22300
188	1/28/98	188	RTA00000188AF.o.05.1	M00003806D:G05	4668
189	1/28/98	189	RTA00000197AF.h.10.1	M00001476B:F10	15554
190	1/28/98	190	RTA00000134A.c.7.1	M00001528A:A01	5175
191	1/28/98	191	RTA00000187AF.p.23.1	M00003748B:F02	39804
192	1/28/98	192	RTA00000185AF.m.7.1	M00001605C:D12	39804
193	1/28/98	193	RTA00000199AF.n.3.1	M00003946D:C11	0
194	1/28/98	194	RTA00000200R.k.01.1	M00004188C:A09	40049
195	1/28/98	195	RTA00000198AF.c.10.1	M00001577B:H02	77149
196	1/28/98	196	RTA00000198F.e.10.1	M00001599B:E09	9727

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
197	1/28/98	197	RTA00000198F.I.12.1	M00001669C:B01	8592
198	1/28/98	198	RTA00000197AR.e.07.1	M00001453D:G12	86969
199	1/28/98	199	RTA00000199R.c.09.1	M00003800A:C09	16824
200	1/28/98	200	RTA00000182AF.f.2.1	M00001469D:D02	4794
201	1/28/98	201	RTA00000198AF.p.18.1	M00003769B:D03	23081
202	1/28/98	202	RTA00000200R.1.17.2	M00004217C:D03	12771
203	1/28/98	203	RTA00000201F.d.09.1	M00004380B:A05	1827
204	1/28/98	204	RTA00000180AR.o.5.2	M00001437D:C04	7848
205	1/28/98	205	RTA00000189AF.g.11.1	M00003858D:F12	0
206	1/28/98	206	RTA00000181AF.o.04.2	M00001457C:C12	22205
207	1/28/98	207	RTA00000199AF.l.19.1	M00003924B:D04	22460
208	1/28/98	208	RTA00000198AF.h.22.1	M00001635C:A03	22366
209	1/28/98	209	RTA00000182AF.c.5.1	M00001464D:F06	6397
210	1/28/98	210	RTA00000189AR.b.12.1	M00003829B:G03	17233
211	1/28/98	211	RTA00000199AF.m.15.1	M00003939A:A02	10101
212	1/28/98	212	RTA00000197AF.j.9.1	M00001494B:C01	13236
213	1/28/98	213	RTA00000200F.o.04.1	M00004260D:C12	12514
214	1/28/98	214	RTA00000200AF.f.22.1	M00004121C:F06	16521
215	1/28/98	215	RTA00000192AR.e.14.3	M00004142A:D08	3300
216	1/28/98	216	RTA00000188AF.g.9.1	M00003774B:B08	4959
217	1/28/98	217	RTA00000198AF.h.3.1	M00001625D:C07	22562
218	1/28/98	218	RTA00000188AF.o.18.1	M00003811D:A12	13678
219	1/28/98	219	RTA00000198AF.m.19.1	M00001680D:D02	40041
220	1/28/98	220	RTA00000200AF.h.01.2	M00004141D:A09	0
221	1/28/98	221	RTA00000189AF.i.17.1	M00003868C:H10	16814
222	1/28/98	222	RTA00000185AF.i.4.1	M00001594A:B12	13942
223	1/28/98	223	RTA00000197F.i.9.1	M00001488D:C10	0
224	1/28/98	224	RTA00000188AF.m.11.1	M00003799A:D09	0
225	1/28/98	225	RTA00000189AF.b.5.1	M00003828A:E04	3784
226	1/28/98	226	RTA00000191AR.o.09.4	M00004096A:G02	0
227	1/28/98	227	RTA00000201R.d.02.2	M00004375A:H01	2599
228	1/28/98	228	RTA00000187AR.h.15.2	M00001679A:A06	6660
229	1/28/98	229	RTA00000198AF.g.3.1	M00001615C:F03	0
230	1/28/98	230	RTA00000185AR.b.18.1	M00001575B:C09	12171
231	1/28/98	231	RTA00000192AF.1.13.2	M00004185C:C03	11443
232	1/28/98	232	RTA00000186AF.j.03.2	M00001638A:E07	0
233	1/28/98	233	RTA00000197AF.1.8.1	M00001511B:C06	39954
234	1/28/98	234	RTA00000191AF.f.8.1	M00004035A:A04	6541
235	1/28/98	235	RTA00000201AF.a.02.1	M00004295B:D02	35362
236	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
237	1/28/98	237	RTA00000197AF.k.10.1	M00001500D:B11	0
238	1/28/98	238	RTA00000187AR.k.12.1	M00001679D:F02	78415
239	1/28/98	239	RTA00000201R.d.02.1	M00004375A:H01	2599
240	1/28/98	240	RTA00000178AF.e.1.1	M00001369A:H12	2664
241	1/28/98	241	RTA00000200AF.I.17.1	M00004217C:D03	12771
242	1/28/98	242	RTA00000198AF.m.17.1	M00001679D:F06	77992
243	1/28/98	243	RTA00000181AF.m.15.3	M00001455D:A11	12081
244	1/28/98	244	RTA00000199F.f.12.2	M00003844C:A08	8131
245	1/28/98	245	RTA00000200AF.k.7.1	M00004193C:G11	0
246	1/28/98	246	RTA00000199AF.l.4.1	M00003911D:B04	4410

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
247	Appln 1/28/98	Appln 247	RTA00000198AF.k.08.1	M00001656C:G08	17436
247 248	1/28/98	247	RTA00000198A1.R.00.11	M00001030C:G00	39814
249	1/28/98	249	RTA00000170R.c.14.17 RTA00000200R.o.03.2	M000013702:C01	22807
250	1/28/98	250	RTA00000200R.0.03.2 RTA00000192AF.j.21.1	M00004237C:F100 M00004176D:B12	2289
250 251	1/28/98	250	RTA00000192AF.n.13.1	M00004170D:B12	8210
	1/28/98	252	RTA00000192AI .ii.13.1 RTA00000181AF.e.18.3	M00004197D:1101 M00001448D:C09	8
252	1/28/98	252 253	RTA00000181AF.e.17.3	M00001448D:C09	8
253		253 254	RTA00000178AF.n.2.1	M00001446D:C07	17083
254	1/28/98			M00001383C:1111 M00003889A:D10	5121
255	1/28/98	255	RTA00000199AF.j.17.1	M00003889A.D10 M00001549C:E06	16347
256	1/28/98	256	RTA00000184AR.e.15.1	M00001549C:E00	9810
257	1/28/98	257	RTA00000198AF.e.20.1	M00001804C.E09 M00003868B:D12	16621
258	1/28/98	258	RTA00000199F.h.12.2		17209
259	1/28/98	259	RTA00000197AF.j.4.1	M00001492D:A11	77992
260	1/28/98	260	RTA00000198R.m.17.1	M00001679D:F06	13183
261	1/28/98	261	RTA00000192AF.a.24.1	M00004114C:F11	8551
262	1/28/98	262	RTA00000186AF.c.17.1	M00001619D:G05	
263	1/28/98	263	RTA00000190AF.n.6.1	M00003965A:B11	0
264	1/28/98	264	RTA00000179AF.k.3.3	M00001401A:H07	0
265	1/28/98	265	RTA00000177AF.e.14.1	M00001343D:H07	23255
266	1/28/98	266	RTA00000199F.f.21.2	M00003847C:E09	13344
267	1/28/98	267	RTA00000186AF.g.11.2	M00001630B:H09	5214
268	1/28/98	268	RTA00000186AF.h.01.2	M00001632A:F12	0
269	1/28/98	269	RTA00000183AF.k.13.1	M00001534B:C12	0
270	1/28/98	270	RTA00000178R.1.08.1	M00001383A:C03	39648
271	1/28/98	271	RTA00000201F.d.02.1	M00004375A:H01	2599
272	1/28/98	272	RTA00000199F.g.08.2	M00003853D:G08	0
273	1/28/98	273	RTA00000201F.c.08.1	M00004353C:H07	0
274	1/28/98	274	RTA00000191AF.o.17.1	M00004102A:H02	5957
275	1/28/98	275	RTA00000191AF.o.17.2	M00004102A:H02	5957
276	1/28/98	276	RTA00000198AF.j.15.1	M00001653B:E09	4369
277	1/28/98	277	RTA00000198AR.i.08.1	M00001639A:F10	9807
278	1/28/98	278	RTA00000198AF.p.16.1	M00003768A:E02	71877
279	1/28/98	279	RTA00000196AF.h.24.1	M00001386A:D11	7308
280	1/28/98	280	RTA00000193AF.b.18.1	M00004233C:H09	7542
281	1/28/98	281	RTA00000188AF.n.10.1	M00003802D:B11	10283
282	1/28/98	282	RTA00000193AF.c.15.1	M00004248B:E08	3726
283	1/28/98	283	RTA00000177AF.i.8.4	M00001350A:H01	7187
284	1/28/98	284	RTA00000199F.d.10.2	M00003808C:B05	22049
285	1/28/98	285	RTA00000181AR.j.14.3	M00001453B:E10	5399
286	1/28/98	286	RTA00000181AR.k.2.3	M00001453C:A11	0
287	1/28/98	287	RTA00000200AF.b.07.1	M00004039C:C01	17125
288	1/28/98	288	RTA00000181AR.i.06.3	M00001452A:C07	19119
289	1/28/98	289	RTA00000196F.k.07.1	M00001399C:D09	22443
290	1/28/98	290	RTA00000201F.f.10.1	M00004498D:D05	5231
291	1/28/98	291	RTA00000200AF.e.16.1	M00004101C:G08	12068
292	1/28/98	292	RTA00000199AF.m.18.1	M00003939C:F04	0
293	1/28/98	293	RTA00000197AF.e.13.1	M00001454C:F02	662
294	1/28/98	294	RTA00000198AF.k.23.1	M00001661B:C08	8995
295	1/28/98	295	RTA00000181AR.i.19.2	M00001452C:B06	16970
296	1/28/98	296	RTA00000196AF.f.20.1	M00001371D:G01	22774

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
297	1/28/98	297	RTA00000178AF.f.9.3	M00001371C:E09	7172
298	1/28/98	298	RTA00000197AR.e.11.1	M00001454B:G03	2306
299	1/28/98	299	RTA00000196AF.f.5.1	M00001366D:G02	11937
300	2/24/98	464	RTA00000195AF.c.12.1	M00003818B:G12	37582
300	1/28/98	300	RTA00000195AF.c.12.1	M00003818B:G12	37582
301	1/28/98	301	RTA00000181AR.i.19.3	M00001452C:B06	16970
302	1/28/98	302	RTA00000186AF.d.1.2	M00001621C:C08	40044
303	1/28/98	303	RTA00000186AR.e.03.3	M00001623D:C10	22110
304	1/28/98	304	RTA00000182AR.c.5.1	M00001464D:F06	6397
305	1/28/98	305	RTA00000200AF.b.15.1	M00004040D:F01	10627
306	1/28/98	306	RTA00000199AF.p.12.1	M00003989A:H11	12578
307	1/28/98	307	RTA00000200F.n.05.2	M00004246C:A09	18989
308	1/28/98	308	RTA00000178AF.j.20.1	M00001380C:E05	15066
309	1/28/98	309	RTA00000198AF.h.12.1	M00001632C:A02	9503
310	1/28/98	310	RTA00000188AF.m.08.1	M00003798D:H08	22155
311	1/28/98	311	RTA00000191AR.j.4.2	M00004071D:A10	5198
312	1/28/98	312	RTA00000193AF.h.2.1	M00004290A:B03	3273
313	1/28/98	313	RTA00000183AF.o.11.1	M00001540D:D02	0
314	1/28/98	314	RTA00000182AF.o.5.1	M00001493B:D09	5007
315	1/28/98	315	RTA00000199R.d.23.1	M00003815D:H09	37477
316	1/28/98	316	RTA00000198AF.h.24.1	M00001636C:C01	8390
317	1/28/98	317	RTA00000198AF.p.09.1	M00003761D:E02	10473
318	1/28/98	318	RTA00000200AF.g.17.1	M00004138A:H09	0
319	1/28/98	319	RTA00000200F.n.05.1	M00004246C:A09	18989
320	1/28/98	320	RTA00000196AF.m.13.1	M00001415B:E09	16290
321	1/28/98	321	RTA00000181AR.b.21.1	M00001444C:D05	3266
322	1/28/98	322	RTA00000184AR.b.21.1	M00001546B:B02	39788
323	1/28/98	323	RTA00000182AF.m.21.1	M00001490C:C12	18699
324	1/28/98	324	RTA00000184F.j.06.1	M00001556B:G02	11294
325	1/28/98	325	RTA00000182AF.d.18.4	M00001467D:H05	37435
326	1/28/98	326	RTA00000197AR.e.19.1	M00001455D:A09	8047
327	1/28/98	327	RTA00000182AF.i.1.3	M00001479B:A01	7033
328	1/28/98	328	RTA00000200AF.g.09.1	M00004131B:H09	22785
329	1/28/98	329	RTA00000186AF.b.9.1	M00001616C:F07	0
330	1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
331	1/28/98	331	RTA00000197AR.c.20.1	M00001449D:A06	16282
332	1/28/98	332	RTA00000193AR.n.04.3	M00004375C:D01	9850
333	1/28/98	333	RTA00000196F.k.15.1	M00001400A:F06	8320
334	1/28/98	334	RTA00000181AR.b.21.3	M00001444C:D05	3266
335	1/28/98	335	RTA00000182AF.e.3.2	M00001468B:H06	0
336	1/28/98	336	RTA00000186AF.f.24.1	M00001629B:E06	0
337	1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
338	1/28/98	338	RTA00000184AF.i.1.1	M00001554B:C07	0
339	1/28/98	339	RTA00000193AF.d.1.1	M00004250D:D10	0
340	1/28/98	340	RTA00000185AF.n.8.1	M00001608B:A03	0
341	1/28/98	341	RTA00000181AF.I.06.2	M00001454C:C08	0
342	1/28/98	342	RTA00000196AF.d.10.1	M00001354C:B06	22256
343	1/28/98	343	RTA00000201F.a.18.1	M00004314B:G07	16837
344	1/28/98	344	RTA00000198AF.o.02.1	M00003748A:B07	68756
345	1/28/98	345	RTA00000187AF.h.21.1	M00001679A:F01	39171

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority Appln	Priority Appln			
346	1/28/98	346	RTA00000197AR.k.22.1	M00001505C:H01	11394
347	1/28/98	347	RTA00000199F.b.03.2	M00003779B:E12	38340
348	1/28/98	348	RTA00000200F.n.07.2	M000037472:212	8663
349	1/28/98	349	RTA00000191AF.j.15.1	M00004073B:B01	6308
350	1/28/98	350	RTA00000193AR.c.7.2	M00004241D:F11	9850
351	1/28/98	351	RTA00000179AF.c.22.1	M00001211B:F11	22515
352	1/28/98	352	RTA00000177AF.p.3.1	M00001550A:A03	7239
353	1/28/98	353	RTA00000198F.a.9.1	M00001557D:C08	0
354	1/28/98	354	RTA00000198R.k.03.1	M00001655A:F06	22765
355	1/28/98	355	RTA00000184AR.b.24.1	M00001546B:C05	5777
356	1/28/98	356	RTA00000180AF.I.12.2	M00001433B:H11	0
357	1/28/98	357	RTA00000184AF.o.15.1	M00001564D:C09	0
358	1/28/98	358	RTA00000198AF.g.7.1	M00001616C:C09	13386
359	1/28/98	359	RTA00000196AF.b.17.1	M00001348A:D04	12193
360	1/28/98	360	RTA00000198F.i.5.1	M00001638A:D10	39989
361	1/28/98	361	RTA00000177AR.g.16.4	M00001337A:B10	13576
362	1/28/98	362	RTA00000197AR.c.24.1	M00001450A:B12	82498
363	1/28/98	363	RTA00000196AF.e.14.1	M00001362C:A10	12850
364	1/28/98	364	RTA00000187AF.g.13.1	M00001676C:C11	2991
365	1/28/98	365	RTA00000196F.I.20.2	M00001410B:G05	22678
366	1/28/98	366	RTA00000192AF.o.19.1	M00004208D:H08	3549
367	1/28/98	367	RTA00000196F.i.24.1	M00001392C:D10	4233
368	1/28/98	368	RTA00000198AF.k.18.1	M00001660A:C12	17432
369	1/28/98	369	RTA00000196F.m.3.1	M00001413A:F02	10453
370	1/28/98	370	RTA00000179AF.c.15.3	M00001392D:H06	2995
371	1/28/98	371	RTA00000197F.e.7.1	M00001453D:G12	86969
372	1/28/98	372	RTA00000186AF.d.23.1	M00001623B:G07	22187
373	1/28/98	373	RTA00000196F.e.12.1	M00001361C:H11	10147
374	1/28/98	374	RTA00000178AF.I.11.1	M00001383A:G04	23286
375	1/28/98	375	RTA00000177AF.m.18.1	M00001355B:G11	0
376	1/28/98	376	RTA00000177AF.m.18.3	M00001355B:G11	0
377	1/28/98	377	RTA00000178AF.m.19.1	M00001384D:H07	0
378	1/28/98	378	RTA00000181AF.k.24.3	M00001454B:C12	7005
379	1/28/98	379	RTA00000180AF.1.06.2	M00001433A:G07	5625
380	1/28/98	380	RTA00000182AF.k.24.1	M00001485D:B10	5625
381	1/28/98	381	RTA00000199AF.m.14.1	M00003938A:B04	10580
382	1/28/98	382	RTA00000200AF.j.6.1	M00004176B:E08	22902
383	1/28/98	383	RTA00000199F.f.20.2	M00003847B:G03	0
384	1/28/98	384	RTA00000196AF.h.17.1	M00001384C:F12	39215
385	1/28/98	385	RTA00000201F.c.24.1	M00004374D:E10	35731
386	1/28/98	386	RTA00000197AR.j.04.1	M00001492D:A11	17209
387	2/24/98	632	RTA00000191AF.j.14.1	M00004073A:H12	1002
387	1/28/98	387	RTA00000191AF.j.14.1	M00004073A:H12	1002
388	1/28/98	388	RTA00000185AF.n.17.1	M00001609B:A11	5336
389	1/28/98	389	RTA00000181AR.k.2.2	M00001453C:A11	0
390	1/28/98	390	RTA00000197AR.f.07.1	M00001457C:C11	19261
391	1/28/98	391	RTA00000179AF.e.20.3	M00001396A:C03	4009
392	1/28/98	392	RTA00000185AF.b.11.2	M00001573C:D03	9024
393	1/28/98	393	RTA00000188AF.b.14.1	M00003754D:D02	0
394	1/28/98	394	RTA00000198AF.p.22.1	M00003771A:G10	0
			141		

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
205	Appln	Appln	PT 4 00000106P = 21 2	M00001252C-1110	Λ
395	1/28/98	395 396	RTA00000196R.c.21.2	M00001352C:H10	0
396	1/28/98		RTA00000179AR.b.02.3	M00001391B:G12	38956
397 398	1/28/98	397 398	RTA00000198AF.b.22.1 RTA00000177AR.l.13.3	M00001571B:E03 M00001353A:G12	38936 8078
398 399	1/28/98	398 399		M00001333A:G12	40122
	1/28/98		RTA00000186AF.m.15.2	M00001649C:B10 M00001623D:G03	
400	1/28/98	400	RTA00000186AR.e.07.3		4175
401	1/28/98	401	RTA00000195F.e.04.1	M00004465B:D04	6731
402	1/28/98	402	RTA00000177AF.b.21.4	M00001341A:F12	4443
403	1/28/98	403	RTA00000184AF.f.13.1	M00001550D:H02	3784
404	1/28/98	404	RTA00000195AF.b.6.1	M00001446C:G10	39490
405	1/28/98	405	RTA00000197AF.b.24.1	M00001446C:D09	23171
406	1/28/98	406	RTA00000199F.f.09.2	M00003842B:D09	22907
407	1/28/98	407	RTA00000178AF.e.20.1	M00001370D:E12	3135
408	1/28/98	408	RTA00000183AR.I.15.1	M00001535C:E01	39383
409	1/28/98	409	RTA00000180AF.d.1.3	M00001418D:B06	8526
410	1/28/98	410	RTA00000201F.a.20.1	M00004316A:G09	22639
411	1/28/98	411	RTA00000179AF.j.13.3	M00001400B:H06	0
412	2/24/98	678	RTA00000195AF.c.24.1	M00003860D:H07	0
412	1/28/98	412	RTA00000195AF.c.24.1	M00003860D:H07	0
413	1/28/98	413	RTA00000200F.a.12.1	M00004031D:B05	16751
414	1/28/98	414	RTA00000185AR.b.15.1	M00001573D:F04	39813
415	1/28/98	415	RTA00000200AF.f.09.1	M00004111C:E11	12863
416	1/28/98	416	RTA00000199F.a.5.1	M00003773B:G01	22134
417	1/28/98	417	RTA00000200R.d.16.1	M00004085A:B02	39875
418	1/28/98	418	RTA00000187AR.k.01.1	M00001679D:B05	78356
419	1/28/98	419	RTA00000182AF.j.20.1	M00001483B:D03	4769
420	1/28/98	420	RTA00000181AF.c.11.1	M00001445D:A06	4769
421	1/28/98	421	RTA00000200AF.i.21.1	M00004167D:A07	5316
422	1/28/98	422	RTA00000189AF.b.12.1	M00003829B:G03	17233
423	1/28/98	423	RTA00000188AR.b.17.1	M00003755A:B03	10662
424	1/28/98	424	RTA00000187AR.j.24.1	M00001679D:B05	78356
425	1/28/98	425	RTA00000200AF.c.16.1	M00004064D:A11	23433
426	1/28/98	426	RTA00000199AF.o.19.1	M00003980D:E09	36927
427	1/28/98	427	RTA00000187AR.d.9.2	M00001664D:G07	5483
428	1/28/98	428	RTA00000185AF.b.15.2	M00001573D:F04	39813
429	1/28/98	429	RTA00000196F.i.19.1	M00001390C:C11	39498
430	1/28/98	430	RTA00000198R.k.23.1	M00001661B:C08	8995
431	1/28/98	431	RTA00000199AF.k.15.1	M00003905C:G10	8275
432	1/28/98	432	RTA00000198AF.o.05.1	M00003750A:D01	26702
433	1/28/98	433	RTA00000198R.j.18.1	M00001653B:G07	22759
434	1/28/98	434	RTA00000187AR.d.2.2	M00001664C:H10	4892
435	1/28/98	435	RTA00000182AR.c.22.1	M00001467A:D08	16283
436	1/28/98	436	RTA00000200AF.k.11.1	M00004197C:F03	9796
437	1/28/98	437	RTA00000198R.a.23.1	M00001563B:D11	10700
438	1/28/98	438	RTA00000180AR.g.03.4	M00001425A:C11	9024
439	1/28/98	439	RTA00000185AF.d.14.2	M00001579D:G07	8071
440	1/28/98	440	RTA00000177AR.f.13.4	M00001345A:G11	10480
441	1/28/98	441	RTA00000185AF.e.6.1	M00001583B:E10	0
442	1/28/98	442	RTA00000191AF.I.9.1	M00004081C:H06	Ö
443	1/28/98	443	RTA00000197AR.i.17.1	M00001490A:E11	3516
-	,	· ·=	142		
			142		

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
4.4.4	Appln	Appln	DTA00000180AE1161	M00003879A:G05	0
444	1/28/98	444	RTA00000189AF.l.16.1 RTA00000196AF.n.13.1	M00003879A.G03 M00001422C:F12	8396
445	1/28/98	445		M00001422C:F12 M00001463A:F06	9755
446	1/28/98	446	RTA00000182AF.a.23.3		0
447	1/28/98	447	RTA00000198AF.d.8.1	M00001587A:H03	8608
448	1/28/98	448	RTA00000200AF.j.9.1	M00004177C:A01	
449	1/28/98	449	RTA00000181AF.m.22.3	M00001455D:F09	9283
450	1/28/98	450	RTA00000181AF.m.21.3	M00001455D:F09	9283
451	1/28/98	451	RTA00000200AF.b.20.1	M00004043A:D02	40403
452	1/28/98	452	RTA00000199F.d.19.2	M00003813D:H12	6707
453	1/28/98	453	RTA00000199AF.i.20.1	M00003881A:D09	9544
454	1/28/98	454	RTA00000200R.d.04.1	M00004078A:A06	5506
455	1/28/98	455	RTA00000198AF.d.12.1	M00001589A:C01	21142
456	1/28/98	456	RTA00000200AF.b.12.1	M00004040B:F10	22053
457	1/28/98	457	RTA00000191AR.I.7.2	M00004081C:D12	14391
458	1/28/98	458	RTA00000199R.d.16.1	M00003812C:A05	24191
459	1/28/98	459	RTA00000179AF.c.22.3	M00001393B:B09	22515
460	1/28/98	460	RTA00000179AF.c.15.1	M00001392D:H06	2995
461	1/28/98	461	RTA00000190AF.e.13.1	M00003908A:H09	38961
462	1/28/98	462	RTA00000196AF.n.17.1	M00001423D:A09	12477
463	1/28/98	463	RTA00000177AR.k.23.4	M00001352D:D02	35550
464	1/28/98	464	RTA00000199AF.I.14.1	M00003917A:D02	22865
465	1/28/98	465	RTA00000187AF.k.20.1	M00001680B:C01	3648
466	1/28/98	466	RTA00000177AF.p.20.1	M00001361A:A05	4141
467	1/28/98	467	RTA00000195AF.b.19.1	M00001589A:D12	77678
468	. 1/28/98	468	RTA00000198AF.a.18.1	M00001561C:E11	0
469	1/28/98	469	RTA00000190AF.n.2.1	M00003963A:E03	5650
470	1/28/98	470	RTA00000198AF.f.16.1	M00001614A:E06	0
471	1/28/98	471	RTA00000188AF.e.2.1	M00003763B:H01	0
472	1/28/98	472	RTA00000192AF.p.17.1	M00004214C:H05	11451
473	1/28/98	473	RTA00000196F.i.3.1	M00001387A:E10	0
474	1/28/98	474	RTA00000192AR.d.1.3	M00004130D:H01	14507
475	1/28/98	475	RTA00000187AR.m.3.3	M00001682C:B12	17055
476	1/28/98	476	RTA00000200R.g.15.1	M00004135B:G01	22898
477	1/28/98	477	RTA00000180AR.e.22.2	M00001423A:G05	7714
478	1/28/98	478	RTA00000192AR.o.24.2	M00004210B:B05	7191
479	1/28/98	479	RTA00000197R.I.22.1	M00001528A:C11	6962
480	1/28/98	480	RTA00000181AF.o.08.2	M00001457C:H12	849
481	1/28/98	481	RTA00000179AR.I.22.2	M00001405B:E09	4314
482	1/28/98	482	RTA00000187AF.j.7.1	M00001679C:F01	78091
483	1/28/98	483	RTA00000192AF.h.19.1	M00004162C:A07	4642
484	1/28/98	484	RTA00000199F.g.20.2	M00003860D:A01	15767
485	1/28/98	485	RTA00000196AF.c.14.1	M00001352B:F04	23105
486	1/28/98	486	RTA00000190AR.p.22.2	M00003979A:E11	16368
487	1/28/98	487	RTA00000198F.i.8.1	M00001639A:F10	9807
488	1/28/98	488	RTA00000179AR.I.22.4	M00001405B:E09	4314
489	1/28/98	489	RTA00000186AF.h.22.1	M00001634B:C10	16485
490	1/28/98	490	RTA00000198AF.n.05.1	M00001687A:G01	24157
491	1/28/98	491	RTA00000196F.k.11.1	M00001399C:H12	3
492	1/28/98	492	RTA00000198AF.b.8.1	M00001567C:H12	22636
493	1/28/98	493	RTA00000177AF.m.17.1	M00001355B:G10	14391

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority		·	
	Appln	Appln	DTA00000000AE	M00004188C:A09	40049
494	1/28/98	494	RTA00000200AF.k.1.1	M00004188C.A09	0
495	1/28/98	495	RTA00000185AF.j.21.1	M00001397A:E12 M00003975B:F03	2378
496	1/28/98	496	RTA00000190AF.p.3.1 RTA00000198AF.o.09.1	M00003973B:103 M00003751B:A05	4310
497	1/28/98	497		M00003731B:A03 M00003917C:D03	12977
498	1/28/98	498	RTA00000190AF.h.12.1	M00003717C:E09	17018
499	1/28/98	499	RTA00000199F.b.22.2 RTA00000179AR.m.07.5	M00003791C:E09	0
500	1/28/98	500	RTA00000179AK.III.07.3	M00001403D:D11 M00004197C:F03	9796
501	1/28/98	501	RTA00000200R.R.11.1 RTA00000197AF.o.23.1	M00004197C:103	12682
502	1/28/98	502	RTA00000197AF.b.23.1 RTA00000197AF.k.9.1	M00001549A:A09	3138
503	1/28/98	503	RTA00000197AF.k.3.1 RTA00000198AF.g.2.1	M00001500C:C00	16640
504	1/28/98	504 505	RTA00000198AF.g.2.1	M00001813C:B02	9443
505	1/28/98	505 506	RTA00000188A1.ii.03.1 RTA00000198R.o.09.1	M00003301B:B10	4310
506	1/28/98	506 507	RTA00000198AF.c.5.1	M00003731D:F10	53802
507	1/28/98	508	RTA000001387AF.i.14.2	M00001575B:H07	19406
508	1/28/98	508 509	RTA00000187AF.1.14.2 RTA00000183AF.p.17.1	M00001677B:H07	5158
509	1/28/98 1/28/98	510	RTA00000178AF.n.23.1	M0000134371:1112 M00001387B:E02	3298
510 511	1/28/98	511	RTA00000176AF.g.10.1	M00001307B:202	12498
512	1/28/98	512	RTA00000191AF.c.3.1	M00003987D:D06	3549
512	1/28/98	513	RTA00000197AF.h.14.1	M00001477B:F04	7045
513	1/28/98	514	RTA00000196AF.n.02.1	M00001417D:A04	39260
515	1/28/98	515	RTA00000196AF.f.18.1	M00001370D:A12	14506
516	1/28/98	516	RTA00000200AF.e.23.1	M00004107B:A06	14686
517	1/28/98	517	RTA00000184AF.e.14.1	M00001549C:D02	16347
518	1/28/98	518	RTA00000199AF.n.22.1	M00003971A:A06	23064
519	1/28/98	519	RTA00000183AF.a.24.2	M00001499B:A11	10539
520	1/28/98	520	RTA00000195AF.c.8.1	M00001678B:H01	0
520	2/24/98	958	RTA00000195AF.c.8.1	M00001678B:H01	0
521	1/28/98	521	RTA00000197AF.p.12.1	M00001552B:G05	0
522	1/28/98	522	RTA00000178AR.h.17.2	M00001376A:C05	23824
523	1/28/98	523	RTA00000198AF.d.4.1	M00001586D:E02	22435
524	1/28/98	524	RTA00000191AF.j.24.1	M00004076B:G03	0
525	1/28/98	525	RTA00000198AF.c.7.1	M00001575D:G05	19181
526	1/28/98	526	RTA00000185AF.e.20.1	M00001585A:D06	5865
527	1/28/98	527	RTA00000198R.m.23.1	M00001684B:G03	38469
528	1/28/98	528	RTA00000200F.n.09.2	M00004249D:B08	12391
529	1/28/98	529	RTA00000178AF.b.13.1	M00001364A:E11	3114
530	1/28/98	530	RTA00000185AF.d.24.2	M00001582D:F05	0
531	1/28/98	531	RTA00000195F.a.3.1	M00001368A:A03	27179
532	1/28/98	532	RTA00000177AF.o.4.1	M00001358C:C06	0
533	1/28/98	533	RTA00000177AR.m.13.4	M00001355A:C12	4175
534	1/28/98	534	RTA00000201AF.e.01.1	M00004405D:C04	11397
535	1/28/98	535	RTA00000196AF.n.19.1	M00001423D:D12	6881 0
536	1/28/98	536	RTA00000193AR.a.2.3	M00004216D:D03	0
537	1/28/98	537	RTA00000188AF.g.14.1	M00003774C:D02	4175
538	1/28/98	538	RTA00000177AR.m.13.3	M00001355A:C12 M00001445B:E04	9560
539	1/28/98	539	RTA00000197AR.b.13.1 RTA00000179AF.b.10.3	M00001391D:D10	9300
540	1/28/98	540 541	RTA00000179AF.b.10.3 RTA00000197AR.b.16.1	M00001391D:D10	0
541 542	1/28/98	541 542	RTA00000197AR.D.10.1 RTA00000198R.p.12.1	M00001443C:A08	8878
542	1/28/98	242	K1700000130K.p.12.1	14100003703D.D10	5576

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appin	Appln			
543	1/28/98	543	RTA00000200AF.i.19.1	M00004167A:H03	14722
544	1/28/98	544	RTA00000196F.j.13.1	M00001396D:B03	23170
545	1/28/98	545	RTA00000196F.a.2.1	M00001338B:E02	3575
546	1/28/98	546	RTA00000197F.i.6.1	M00001487C:D06	12149
547	1/28/98	547	RTA00000196AF.g.8.1	M00001375B:G12	39665
548	1/28/98	548	RTA00000179AF.f.23.3	M00001397B:G03	35258
549	1/28/98	549	RTA00000198AF.c.16.1	M00001579C:B11	26801
550	1/28/98	550	RTA00000183AF.g.14.1	M00001513D:A03	0
551	1/28/98	551	RTA00000200AR.c.24.1	M00004076D:D04	15972
552	1/28/98	552	RTA00000193AF.b.24.1	M00004237D:D08	35
553	1/28/98	553	RTA00000201F.b.22.1	M00004344B:H04	35728
554	1/28/98	554	RTA00000186AR.e.07.4	M00001623D:G03	4175
555	1/28/98	555	RTA00000198AF.j.08.1	M00001651B:A11	10983
556	1/28/98	556	RTA00000199F.f.17.2	M00003845D:B04	22905
557	1/28/98	557	RTA00000198AF.d.9.1	M00001587D:A10	8841
558	1/28/98	558	RTA00000186AR.h.14.1	M00001632D:H07	0
559	1/28/98	559	RTA00000197AF.p.20.1	M00001554B:B07	22795
560	1/28/98	560	RTA00000184AF.i.23.3	M00001556A:F11	1577
561	1/28/98	561	RTA00000185AR.d.10.1	M00001579C:H10	0
562	1/28/98	562	RTA00000196F.j.12.1	M00001396A:H03	19294
563	1/28/98	563	RTA00000192AR.o.16.2	M0000139071.1103 M00004208B:F05	9061
564	1/28/98	564	RTA000001927IR:0:10.2	M00004138B:B11	1600
565	1/28/98	565	RTA00000191AF.c.10.1	M00004130B:B11	40422
566	1/28/98	566	RTA00000195F.a.4.1	M00003303E:11	20470
567	1/28/98	567	RTA00000177AR.m.13.1	M000013726:G12	4175
568	1/28/98	568	RTA00000196AF.p.01.2	M0000133071:012	87143
569	1/28/98	569	RTA00000196AF.I.23.1	M00001430A:A02	12052
570	1/28/98	570	RTA00000183AF.a.19.2	M00001499A:A05	3788
571	1/28/98	571	RTA00000198AF.b.14.1	M00001569C:B06	801
572	1/28/98	572	RTA00000181AF.I.16.2	M00001454D:E05	13532
573	1/28/98	573	RTA00000196AF.b.7.1	M00001344A:G07	7774
574	1/28/98	574	RTA00000192AF.f.3.1	M00004146C:C11	5257
575	1/28/98	575	RTA00000186AF.I.12.2	M00001645A:C12	19267
576	1/28/98	576	RTA00000196AF.c.7.1	M00001350B:G11	0
577	1/28/98	577	RTA00000190AF.a.24.2	M00003901B:A05	0
578	1/28/98	578	RTA00000180AF.g.17.1	M00001426A:A09	16653
579	1/28/98	579	RTA00000200F.i.7.1	M00004157D:B03	22322
580	1/28/98	580	RTA00000197F.a.12.1	M00001438B:B09	7895
581	1/28/98	581	RTA00000191AF.p.3.2	M00004104B:F11	17
582	1/28/98	582	RTA00000178AR.d.12.4	M00001368A:D07	2476
583	1/28/98	583	RTA00000190AR.h.12.2	M00003917C:D03	12977
584	1/28/98	584	RTA00000190AR.c.03.1	M00003904C:A08	0
585	1/28/98	585	RTA00000198AF.n.18.1	M00001771A:A07	16715
586	1/28/98	586	RTA00000199R.o.11.1	M00003976C:A10	23172
587	1/28/98	587	RTA00000199F.a.3.1	M00003772D:E10	16617
588	1/28/98	588	RTA00000191AF.b.4.1	M00003983C:F03	14936
589	1/28/98	589	RTA00000192AF.I.1.1	M00004183C:D07	16392
590	1/28/98	590	RTA00000190AF.d.2.1	M00003906B:F12	2444
591	1/28/98	591	RTA00000197AF.h.1.1	M00001470A:H01	13075
592	1/28/98	592	RTA00000186AF.e.18.1	M00001624C:A06	0

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
593	1/28/98	593	RTA00000196R.c.14.2	M00001352B:F04	23105
594	1/28/98	594	RTA00000181AR.e.04.3	M00001448A:G09	11825
595	1/28/98	595	RTA00000195R.a.06.1	M00001394A:E04	35265
595	2/24/98	1065	RTA00000195R.a.06.1	M00001394A:E04	35265
596	1/28/98	596	RTA00000184AF.d.9.1	M00001548A:B11	6515
597	1/28/98	597	RTA00000198F.a.4.1	M00001557A:F01	9635
598	1/28/98	598	RTA00000197F.e.10.1	M00001454B:D08	13154
599	1/28/98	599	RTA00000179AF.o.5.1	M00001408D:D04	6172
600	1/28/98	600	RTA00000177AF.g.4.1	M00001346B:B07	4119
601	1/28/98	601	RTA00000184AF.i.10.2	M00001555A:B01	3744
602	1/28/98	602	RTA00000195AF.b.21.1	M00001595B:A09	39055
602	2/24/98	317	RTA00000195AF.b.21.1	M00001595B:A09	39055
603	1/28/98	603	RTA00000133AR.d.11.3	M00001504D:G06	6420
604	1/28/98	604	RTA00000103AR.d.11.5 RTA00000200AF.j.15.1	M00001301B:800	5849
605	1/28/98	605	RTA00000200A1.J.15.1	M00004103D:B01	23300
606	1/28/98	606	RTA000001701.c.5.1 RTA00000179AR.e.01.4	M0000130174:1107 M00001395A:C09	2493
607	1/28/98	607	RTA00000179AR.c.01.4 RTA00000200AF.k.12.1	M00001393A:C09	7359
608	1/28/98	608	RTA00000200AF.R.12.1 RTA00000192AF.p.8.1	M00004170B:D02 M00004212B:C07	2379
609	1/28/98	609	RTA00000192AF.p.8.1 RTA00000196AF.n.05.1	M00004212B:C07 M00001418B:F07	12531
610	1/28/98	610	RTA00000190AF.k.2.1	M00001416D:107	35924
	1/28/98	611	RTA00000200AF.R.2.1 RTA00000196F.l.13.2	M00004188D:G08	0
611		612	RTA00000190F.1.13.2 RTA00000197AR.e.22.1	M00001406A:H02	78758
612	1/28/98	613	RTA00000197AR.e.22.1 RTA00000177AF.k.18.4	M00001450A:1102 M00001352C:A05	53729
613	1/28/98 1/28/98	614	RTA00000177AF.R.18.4 RTA00000201F.f.03.1	M00001332C:A03	22633
614 615	1/28/98	615	RTA000002017.1.03.1 RTA00000197R.p.20.1	M00004493B:B07	22795
616	1/28/98	616	RTA00000197R.p.20.1 RTA00000188AF.m.07.1	M00001334B:B07 M00003798D:E03	23183
617	1/28/98	617	RTA00000179AF.d.13.3	M00003798D:E03	6583
618	1/28/98	618	RTA00000179AF.a.14.1	M00001334A:101 M00004111D:A08	6874
619	1/28/98	619	RTA00000192AF.a.14.1 RTA00000201F.g.08.1	M00004111D:A00 M00004692A:E07	0074
620	1/28/98	620	RTA000002011.g.08.1	M00004692A:E07	0
621	1/28/98	621	RTA00000201R.g.08.1 RTA00000201R.g.08.2	M00004692A:E07	0
622	1/28/98	622	RTA00000201R.g.00.2 RTA00000186AR.m.14.2	M00004072A:E07	9800
623	1/28/98	623	RTA00000180AK.iii.14.2	M00001049B:G12	19047
	1/28/98	624	RTA00000198R.b.24.1 RTA00000200F.o.15.1	M00001371B:B11	7866
624 625	1/28/98	625	RTA000002001.0.13.11 RTA00000196AF.c.19.1	M00004273A:B03	5935
626	1/28/98	626	RTA00000190A1.c.13.11	M00001532C:G03	6539
627	1/28/98	627	RTA00000199F.h.15.2	M00003870A:C05	22269
628	1/28/98	628	RTA000001991.ii.13.2 RTA00000198AF.g.16.1	M0000367011:C03	6602
629	1/28/98	629	RTA00000199R.m.23.1	M00003945A:E09	40166
630	1/28/98	630	RTA00000133AR.g.03.2	M00003513R:209	3956
631	1/28/98	631	RTA00000103AR.g.03.2	M00001312D:G03	0
632	1/28/98	632	RTA00000183AR.g.03.1	M00001131D:E03	3956
633	1/28/98	633	RTA00000197F.i.8.1	M00001312D:003	6292
634	1/28/98	634	RTA00000192AF.j.6.1	M00004172C:D08	11494
635	1/28/98	635	RTA00000132A1.j.0.1 RTA00000181AF.p.7.3	M00004172C:D00	38773
636	1/28/98	636	RTA00000196F.k.20.1	M00001400R:E01	6324
637	1/28/98	637	RTA000001701.R.20.1 RTA00000200AF.g.15.1	M00001402B:112	22898
638	1/28/98	638	RTA00000193AF.1.05.2	M00004133B:301 M00004348A:A02	2815
639	1/28/98	639	RTA00000199AF.j.1.1	M00003881C:G09	6006
640	1/28/98	640	RTA00000199AF.f.5.1	M00003909A:H04	5015
	20, 70		·=-=		

641 NT28/98 641 RTA00000198Fa.i.0.1 M00001558A:E11 6695 642 1/28/98 642 RTA00000189AF.i.14.1 M00003868B:G11 0 643 1/28/98 643 RTA00000197F.i.12.1 M00001486C:G10 16245 644 1/28/98 644 RTA00000197F.i.12.1 M00001489B:A06 3605 645 1/28/98 645 RTA00000197F.i.12.1 M00001352A:E02 16245 646 1/28/98 646 RTA00000197F.i.1.1 M00001530B:D10 16488 647 1/28/98 647 RTA00000199AR.i.19.2 M0000386C:B40 7 649 1/28/98 649 RTA00000189AR.i.19.2 M0000384C:B11 88204 650 1/28/98 651 RTA00000189AR.i.22.2 M00001385A:F12 7861 651 1/28/98 652 RTA00000178AR.m.21.5 M00001385A:F12 7861 653 1/28/98 653 RTA00000178AR.in.22.3 M00001376B:A08 19230 657 1/28/98 656 RTA000	SEQ ID NO:	Filing Date of Priority Appln	SEQ ID NO: in Priority Appln	Sequence Name	Clone Name	Cluster ID
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671 1/28/98 671 RTA00000185AR.d.08.1 M00001579C:E09 6562 672 1/28/98 672 RTA00000196AF.h.09.1 M00001382B:F12 8015 673 1/28/98 673 RTA00000199F.m.3.1 M00003931B:A11 0 674 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 675 1/28/98 675 RTA00000179AR.b.21.3 M00001392C:D05 4366 676 1/28/98 676 RTA00000197AR.m.14.1 M00001531B:E09 14879 677 1/28/98 677 RTA00000197AF.i.19.1 M00001490B:H11 39554 678 1/28/98 678 RTA00000197AF.i.19.1 M00001490B:H11 39554 679 1/28/98 679 RTA00000197AF.d.11.1 M00001451C:E01 27260 680 1/28/98 680 RTA00000177AF.f.10.1 M00001451C:E01 27260 681 1/28/98 681 RTA00000180AF.l.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 684 RTA00000200AR.b.11.1 M0000440A:G12 12043 685 1/28/98 685 RTA00000200F.i.9.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 688 RTA00000197AF.g.4.1 M00001452B:C06 4121						
672 1/28/98 672 RTA00000196AF.h.09.1 M00001382B:F12 8015 673 1/28/98 673 RTA00000199F.m.3.1 M00003931B:A11 0 674 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 675 1/28/98 675 RTA00000179AR.b.21.3 M00001392C:D05 4366 676 1/28/98 676 RTA00000197AR.m.14.1 M00001531B:E09 14879 677 1/28/98 677 RTA00000197AF.i.19.1 M00001490B:H11 39554 678 1/28/98 678 RTA00000197AF.d.11.1 M00001490B:H11 39554 679 1/28/98 679 RTA00000197AF.d.11.1 M00001451C:E01 27260 680 1/28/98 680 RTA0000017AF.f.10.1 M00001345A:E01 6420 681 1/28/98 681 RTA00000180AF.l.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 685 <				-		
673 1/28/98 673 RTA00000199F.m.3.1 M00003931B:A11 0 674 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 675 1/28/98 675 RTA00000179AR.b.21.3 M00001392C:D05 4366 676 1/28/98 676 RTA00000197AR.m.14.1 M00001531B:E09 14879 677 1/28/98 677 RTA00000197AF.i.19.1 M00001490B:H11 39554 678 1/28/98 678 RTA00000197AF.i.19.1 M00001490B:H11 39554 679 1/28/98 679 RTA00000197AF.d.11.1 M00001451C:E01 27260 680 1/28/98 680 RTA00000177AF.f.10.1 M00001345A:E01 6420 681 1/28/98 681 RTA00000180AF.l.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 684 RTA00000187AF.j.01.1 M00004040A:G12 12043 685 1/28/98 685	672					
674 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 675 1/28/98 675 RTA00000179AR.b.21.3 M00001392C:D05 4366 676 1/28/98 676 RTA00000197AR.m.14.1 M00001531B:E09 14879 677 1/28/98 677 RTA00000197AF.i.19.1 M00001490B:H11 39554 678 1/28/98 678 RTA00000190AF.j.3.1 M00003922A:D02 2705 679 1/28/98 679 RTA00000197AF.d.11.1 M00001451C:E01 27260 680 1/28/98 680 RTA00000177AF.f.10.1 M00001345A:E01 6420 681 1/28/98 681 RTA00000180AF.I.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 685 RTA0000020AR.b.11.1 M00004040A:G12 12043 685 1/28/98 686 <						
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676 1/28/98 676 RTA00000197AR.m.14.1 M00001531B:E09 14879 677 1/28/98 677 RTA00000197AF.i.19.1 M00001490B:H11 39554 678 1/28/98 678 RTA00000190AF.j.3.1 M00003922A:D02 2705 679 1/28/98 679 RTA00000197AF.d.11.1 M00001451C:E01 27260 680 1/28/98 680 RTA00000177AF.f.10.1 M00001345A:E01 6420 681 1/28/98 681 RTA00000180AF.l.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 684 RTA00000200F.i.9.1 M0000400A:G12 12043 685 1/28/98 685 RTA00000200F.i.9.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 689 RT	675	1/28/98	675			
677 1/28/98 677 RTA00000197AF.i.19.1 M00001490B:H11 39554 678 1/28/98 678 RTA00000190AF.j.3.1 M00003922A:D02 2705 679 1/28/98 679 RTA00000197AF.d.11.1 M00001451C:E01 27260 680 1/28/98 680 RTA00000177AF.f.10.1 M00001345A:E01 6420 681 1/28/98 681 RTA00000180AF.I.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 684 RTA00000200AR.b.11.1 M00004040A:G12 12043 685 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 688 RTA00000197AF.g.4.1 M0000464B:B03 8821 688 1/28/98 689 R	676	1/28/98				
678 1/28/98 678 RTA00000190AF.j.3.1 M00003922A:D02 2705 679 1/28/98 679 RTA00000197AF.d.11.1 M00001451C:E01 27260 680 1/28/98 680 RTA00000177AF.f.10.1 M00001345A:E01 6420 681 1/28/98 681 RTA00000180AF.I.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 684 RTA00000200AR.b.11.1 M00004040A:G12 12043 685 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 688 RTA00000197AF.g.4.1 M0000464B:B03 8821 688 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121	677	1/28/98				
679 1/28/98 679 RTA00000197AF.d.11.1 M00001451C:E01 27260 680 1/28/98 680 RTA00000177AF.f.10.1 M00001345A:E01 6420 681 1/28/98 681 RTA00000180AF.I.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 684 RTA00000200AR.b.11.1 M00004040A:G12 12043 685 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M0000464B:B03 8821 688 1/28/98 689 RTA00000193AF.g.3.1 M0000450D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121	678	1/28/98	678	RTA00000190AF.j.3.1		
680 1/28/98 680 RTA00000177AF.f.10.1 M00001345A:E01 6420 681 1/28/98 681 RTA00000180AF.l.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 684 RTA00000200AR.b.11.1 M00004040A:G12 12043 685 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 689 RTA00000193AF.g.3.1 M00004050D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121	679	1/28/98	679	_		
681 1/28/98 681 RTA00000180AF.I.04.2 M00001432D:F05 11159 682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 684 RTA00000200AR.b.11.1 M00004040A:G12 12043 685 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 689 RTA00000193AF.g.3.1 M00004050D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121	680	1/28/98	680	RTA00000177AF.f.10.1		
682 1/28/98 682 RTA00000125A.j.16.1 M00001544A:E06 0 683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 684 RTA00000200AR.b.11.1 M00004040A:G12 12043 685 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 688 RTA00000193AF.g.3.1 M00004050D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121	681	1/28/98	681	RTA00000180AF.I.04.2		
683 1/28/98 683 RTA00000187AR.j.01.1 M00001679C:D01 79028 684 1/28/98 684 RTA00000200AR.b.11.1 M00004040A:G12 12043 685 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 688 RTA00000193AF.g.3.1 M00004050D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121	682	1/28/98	682	RTA00000125A.j.16.1		
685 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 688 RTA00000193AF.g.3.1 M00004050D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121		1/28/98	683	RTA00000187AR.j.01.1	M00001679C:D01	79028
685 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 688 RTA00000193AF.g.3.1 M00004050D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121		1/28/98	684	RTA00000200AR.b.11.1		
686 1/28/98 686 RTA00000201F.f.07.1 M00004497A:H03 51116 687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 688 RTA00000193AF.g.3.1 M00004050D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121		1/28/98	685	RTA00000200F.i.9.1	M00004159C:F09	
687 1/28/98 687 RTA00000197AF.g.4.1 M00001464B:B03 8821 688 1/28/98 688 RTA00000193AF.g.3.1 M00004050D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121		1/28/98		RTA00000201F.f.07.1	M00004497A:H03	
688 1/28/98 688 RTA00000193AF.g.3.1 M00004050D:A06 5567 689 1/28/98 689 RTA00000197AF.o.4.1 M00001542B:C06 4121				<u> </u>		
1121				RTA00000193AF.g.3.1	M00004050D:A06	
690 1/28/98 690 RTA00000198R.I.21.1 M00001673A:A04 19194						4121
	690	1/28/98	690	RTA00000198R.I.21.1	M00001673A:A04	19194

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
(0.1	Appln	Appln	P.T. 100000105E 10.1		6000
691	1/28/98	691	RTA00000195F.a.10.1	M00001401C:H03	6803
692	1/28/98	692	RTA00000199F.e.4.1	M00003820B:C05	0
693	1/28/98	693	RTA00000198F.m.12.1	M00001679C:D05	4
694	1/28/98	694	RTA00000201R.c.19.1	M00004370A:G05	22357
695	1/28/98	695	RTA00000197F.m.5.1	M00001528C:H04	10872
696	1/28/98	696	RTA00000180AR.d.16.3	M00001419D:C10	11393
697	1/28/98	697	RTA00000193AF.e.21.1	M00004271B:B06	0
698	1/28/98	698	RTA00000179AF.g.1.3	M00001397C:A10	7588
699	1/28/98	699	RTA00000178AF.a.12.1	M00001362B:H06	0
700	1/28/98	700	RTA00000183AF.i.18.2	M00001529D:H02	40129
701	1/28/98	70 1	RTA00000199AF.o.10.1	M00003974C:E04	0
702	1/28/98	702	RTA00000177AR.b.8.5	M00001340B:A06	17062
703	1/28/98	703	RTA00000198F.l.09.1	M00001664B:D06	3611
704	1/28/98	704	RTA00000190AF.o.12.1	M00003972D:C09	3438
705	1/28/98	705	RTA00000196F.i.5.1	M00001387B:A06	0
706	1/28/98	706	RTA00000177AF.i.6.4	M00001350A:B08	0
707	1/28/98	707	RTA00000179AF.p.15.1	M00001411D:F05	5622
708	1/28/98	708	RTA00000201F.f.06.1	M00004496C:H03	23771
709	1/28/98	709	RTA00000192AF.d.18.1	M00004135D:G02	0
710	1/28/98	710	RTA00000196AF.1.3.1	M00001405B:D07	20864
711	1/28/98	711	RTA00000198F.i.2.1	M00001637B:E07	8076
712	1/28/98	712	RTA00000201F.b.21.1	M00004341B:G03	9071
713	1/28/98	713	RTA00000198AF.g.21.1	M00001624A:F09	6273
714	1/28/98	714	RTA00000199R.g.07.1	M00003853D:D03	0
715	1/28/98	715	RTA00000197AR.k.11.1	M00001500D:E10	53758
716	1/28/98	716	RTA00000200F.p.05.1	M00004285C:A08	3984
717	1/28/98	717	RTA00000200F.o.10.2	M00004269B:C08	36432
718	1/28/98	718	RTA00000196F.I.14.2	M00001408B:G06	23144
719	1/28/98	719	RTA00000183AF.b.12.1	M00001500A:B02	0
720	1/28/98	720	RTA00000197AF.f.14.1	M00001459B:C09	3732
721	1/28/98	721	RTA00000180AF.c.4.1	M00001417B:C04	5415
722	1/28/98	722	RTA00000199R.j.24.1	M00003895C:A10	0
723	1/28/98	723	RTA00000183AF.p.24.1	M00001543C:F01	3116
724	1/28/98	724	RTA00000177AR.f.15.4	M00001345B:E10	9062
725	1/28/98	725	RTA00000197AF.b.1.1	M00001343D:E10	12134
726	1/28/98	726	RTA00000200R.f.10.1	M00004111D:B07	4
727	1/28/98	727	RTA00000184AF.n.12.2	M00001561D:C11	3727
728	1/28/98	728	RTA00000177AR.f.17.4	M00001345C:B01	8594
729	1/28/98	729	RTA00000184AF.a.19.1	M00001544C:C06	2628
730	1/28/98	730	RTA00000192AF.o.11.1	M00004205D:F06	0
731	1/28/98	731	RTA00000184F.k.02.1	M00001203B:H00	5192
732	1/28/98	732	RTA00000186AF.p.01.2	M00001654D:G11	40440
733	1/28/98	733	RTA00000200AF.d.20.1	M00004087A:G08	26600
734	1/28/98	734	RTA00000200AF.d.21.1	M00004087C:D03	0
735	1/28/98	735	RTA00000192AF.b.11.1	M00004117A:G01	40014
736	1/28/98	736	RTA00000192AF.o.13.1	M00004117A:G01 M00001428B:A09	0
737	1/28/98	737	RTA00000189AR.m.9.1	M00001420B:A09	2917
738	1/28/98	738	RTA00000183AF.o.8.1	M00003600B:C00	8927
739	1/28/98	739	RTA00000181AF.p.12.3	M00001340C:B10	22204
740	1/28/98	740	RTA00000198AF.d.15.1	M00001400C:H02	5997
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SEQ ID NO:	Filing Date of	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Priority Appln	Appln			
741	1/28/98	741	RTA00000196AF.n.22.1	M00001424B:H04	9572
741	1/28/98	741	RTA00000170AF.m.1.1	M00001424D:1104 M00001353D:D10	14929
742	1/28/98	743	RTA0000017771	M00001333B:F06	16342
743 744	1/28/98	743 744	RTA000001767ti .k.5.1	M00001301D:100	7958
745	1/28/98	745	RTA000001701.iii.4.1 RTA00000183AF.m.11.1	M0000147571.703 M00001536D:G02	8927
743 746	1/28/98	74 <i>5</i> 746	RTA00000178AF.i.01.2	M00001336B:F03	4
740 747	1/28/98	740 747	RTA00000176AF.1.01.2 RTA00000190AF.c.6.1	M00001370D:103	4780
748	1/28/98	748	RTA00000190AF.b.24.1	M00003704D:B10	19047
74 6 749	1/28/98	748 749	RTA00000178AR.i.13.4	M00001377B:H01	0
749 750	1/28/98	749 750	RTA00000178AK.i.13.4 RTA00000198AF.a.19.1	M00001577B:H01	0
750 751	1/28/98	750 751	RTA00000179AF.a.17.1 RTA00000179AF.c.4.3	M00001301D:C03	0
751 752	1/28/98	751 752	RTA00000179AF.c.4.3 RTA00000192AF.o.7.1	M00001392D:B11 M00004204D:C03	5275
752 753	1/28/98	752 753	RTA00000192AF.o.17.1	M00004204D:C03	5275
753 754	1/28/98	753 754	RTA00000192AF.0.17.1 RTA00000187AF.1.11.1	M00004208D:B10	4482
754 755		754 755	RTA00000187AF.1.11.1 RTA00000199F.c.21.2	M00001081A.1 03 M00003803C:D09	5070
	1/28/98		RTA00000199F.C.21.2 RTA00000404F.a.02.1	M00003803C.D09 M00001589B:E12	9738
756	2/24/98	1	RTA00000404F.a.02.1 RTA00000406F.d.16.1	M00001389B.E12 M00003875C:G02	15040
757 758	2/24/98	2		M00003875C:G02 M00004105C:B05	63074
758 750	2/24/98	3 4	RTA00000420F.d.18.1 RTA00000339F.i.20.1	M00004103C:B03 M00001438D:C06	4356
759 760	2/24/98	5	RTA00000339F.1.20.1 RTA00000408F.o.12.2	M00001438D:C00 M00001572A:A10	78578
760	2/24/98				79623
761 762	2/24/98	6	RTA00000119A.j.15.1	M00001460A:E11	19623 66467
762 763	2/24/98	7	RTA00000413F.d.12.1	M00004088C:A12	
763	2/24/98	8 9	RTA00000423F.i.12.1	M00003914D:E03 M00003918C:H10	9118 15051
764 765	2/24/98		RTA00000406F.n.02.1		
765 766	2/24/98	10 11	RTA00000350R.c.12.1 RTA00000411F.k.05.1	M00001550D:A04 M00003850D:B05	9728 64777
766 767	2/24/98	12		M00003830D.B03 M00001366D:E12	10020
767 768	2/24/98	12	RTA00000339F.b.17.1 RTA00000406F.f.18.1	M00001300D:E12 M00003879B:G02	38587
768 769	2/24/98 2/24/98	13	RTA00000419F.b.09.1	M00003879B:G02 M00001694C:F12	78128
7 09 770	2/24/98	15	RTA00000419F.c.19.1	M00001694C.F12 M00003820A:A08	64346
770 771	2/24/98	16	RTA00000419F.c.19.1 RTA00000399F.a.02.1	M00003820A:A08 M00001366D:C12	04340
772	2/24/98	17	RTA000003331.a.02.1 RTA00000411F.m.15.1	M00001300D:C12	78014
773	2/24/98	18	RTA000004111.III.13.1 RTA00000420F.g.12.1	M00003808D:B09	0
773 774	2/24/98	19	RTA000004201.g.12.1 RTA00000123A.k.23.1	M00004873B:G04 M00001533A:G05	80313
775	2/24/98	20	RTA00000123A.R.23.1 RTA00000404F.m.04.2	M00001555A:G05	22720
776	2/24/98	21	RTA00000411F.g.08.1	M00001041A:A11	45815
777	2/24/98	22	RTA00000130A.m.15.1	M00003622B:B04	81630
778	2/24/98	23	RTA00000130A.iii.13.1 RTA00000411F.k.20.1	M00001022A:1112 M00003854B:A07	64973
779	2/24/98	24	RTA00000423F.1.09.1	M00003434B:H07	9752
780	2/24/98	25	RTA00000418F.k.05.1	M0000111071.1100	73021
7 8 1	2/24/98	26	RTA00000423F.h.18.1	M00003876C:D02	37972
782	2/24/98	27	RTA00000420F.n.19.2	M00005259B:C01	0
78 3	2/24/98	28	RTA00000422F.p.06.2	M00001661A:B11	39282
78 4	2/24/98	29	RTA00000404F.n.16.2	M00001649C:D05	39095
785	2/24/98	30	RTA00000411F.m.24.1	M00003870B:B08	77568
7 8 6	2/24/98	31	RTA00000134A.j.10.1	M00001534A:G06	81383
7 8 7	2/24/98	32	RTA00000409F.j.02.1	M00001611B:E06	76417
788	2/24/98	33	RTA00000403F.j.15.1	M00001539B:G07	23840
789	2/24/98	34	RTA00000411F.n.11.1	M00003875A:B01	77276
790	2/24/98	35	RTA00000339F.i.13.1	M00001434A:B10	5970

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
5 0.1	Appln	Appln	D.T. 1000004145 00 4	1400005 0 5 0 5	
791	2/24/98	36	RTA00000414F.e.22.1	M00005257D:A06	0
792	2/24/98	37	RTA00000406F.o.15.1	M00003988D:A08	37482
793	2/24/98	38	RTA00000412F.g.04.2	M00003971B:B07	64457
794	2/24/98	39	RTA00000187AF.I.7.1	M00001680D:F08	10539
795	2/24/98	40	RTA00000352R.I.06.1	M00004187D:H06	40343
796	2/24/98	41	RTA00000419F.b.12.1	M00003806B:C09	63148
797	2/24/98	42	RTA00000423F.k.17.2	M00004038A:F02	37512
798	2/24/98	43	RTA00000420F.g.04.1	M00004891B:B12	0
799	2/24/98	44	RTA00000418F.k.14.1	M00001639A:H06	76133
800	2/24/98	45	RTA00000409F.I.12.1	M00001615A:D06	26755
801	2/24/98	46	RTA00000404F.c.20.1	M00001594A:D08	39088
802	2/24/98	47	RTA00000423F.g.09.1	M00003904C:B06	38958
803	2/24/98	48	RTA00000411F.b.24.1	M00001677B:A12	30041
804	2/24/98	49	RTA00000406F.d.12.1	M00003875C:A01	38575
805	2/24/98	50	RTA00000411F.f.02.1	M00003813A:D08	63386
806	2/24/98	51	RTA00000129A.n.21.1	M00001604A:C11	79381
807	2/24/98	52	RTA00000409F.m.12.1	M00001618B:D09	73490
808	2/24/98	53	RTA00000410F.c.04.1	M00001633D:G09	74099
809	2/24/98	54	RTA00000399F.o.01.1	M00001595C:E01	3055
810	2/24/98	55	RTA00000406F.m.09.1	M00003914C:H05	26891
811	2/24/98	56	RTA00000411F.b.06.1	M00001676C:A04	77884
812	2/24/98	57	RTA00000409F.I.21.1	M00001615B:G07	73143
813	2/24/98	58	RTA00000420F.m.18.1	M00005254D:A10	0
814	2/24/98	59	RTA00000346F.j.08.1	M00003879B:A06	39951
815	2/24/98	60	RTA00000413F.p.17.2	M00005136D:G06	0
816	2/24/98	61	RTA00000410F.n.07.1	M00001662A:G01	78823
817	2/24/98	62	RTA00000339F.n.10.1	M00001453B:F08	13719
818	2/24/98	63	RTA00000404F.1.20.2	M00001639B:H05	38638
819	2/24/98	64	RTA00000413F.d.18.1	M00004090B:B04	65305
820	2/24/98	65	RTA00000404F.p.04.2	M00001652D:E05	39069
821	2/24/98	66	RTA00000405F.g.19.2	M00001632B:E03	37150
822	2/24/98	67	RTA00000409F.a.22.1	M000015731:000 M00001583B:F02	75200
823	2/24/98	68	RTA00000339F.n.03.1	M00001303B:102	0
824	2/24/98	69	RTA00000405F.o.18.1	M00003839A:D07	11016
825	2/24/98	70	RTA00000409F.m.13.1	M00001618B:E05	0
826	2/24/98	71	RTA00000120A.d.24.1	M00001464A:E10	5085
827	2/24/98	72	RTA00000347F.a.08.1	M00001592C:G04	3135
828	2/24/98	73	RTA00000413F.p.15.2	M00005136D:D06	0
829	2/24/98	74	RTA00000408F.e.22.2	M00003130B:B00	26930
830	2/24/98	75	RTA00000350R.i.22.1	M00001470B:100	0
831	2/24/98	76	RTA00000413F.d.16.1	M00001000D:703	63331
832	2/24/98	77	RTA00000420F.j.22.1	M00005173B:F01	0
833	1/28/98	59	RTA00000195AF.b.13.1	M00003173D:101 M00001560D:A03	12605
833	2/24/98	78	RTA00000195AF.b.13.1	M00001560D:A03	12605
834	2/24/98	79 79	RTA00000419F.g.08.1	M00001300D://03	66700
835	2/24/98	80	RTA000004171.g.06.1 RTA00000122A.g.16.1	M00003842C:D11	81366
836	2/24/98	81	RTA00000122A.g.10.1 RTA00000419F.c.16.1	M00001314A:B04 M00003819D:B01	65254
837	2/24/98	82	RTA00000417F.b.03.1	M00003619B:B01	23634
838	2/24/98	83	RTA000004111.b.03.1 RTA00000405F.e.11.2	M00001676B:E01	9331
839	2/24/98	84	RTA000004031.c.11.2 RTA00000352R.i.15.1	M00001003D:C00	4363
337	2127170	דט	K1700000332K.I.13.1	COO. CC 1 FOOD 111	COCF

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
0.40	Appln	Appln	P.T. 4.00000230T1 22 1	14000014070 DOI	5554
840	2/24/98	85	RTA00000339F.k.22.1	M00001427C:D01	5556
841	2/24/98	86	RTA00000346F.g.22.1	M00003794D:G03	6371
842	2/24/98	87	RTA00000403F.1.20.1	M00001573A:A06	18267
843	2/24/98	88	RTA00000420F.i.24.1	M00005134B:E08	0
844	2/24/98	89	RTA00000406F.c.08.1	M00003870C:A10	22387
845	2/24/98	90	RTA00000411F.a.02.1	M00001675B:E02	78537
846	2/24/98	91	RTA00000355R.e.15.1	M00004316A:G09	22639
847	2/24/98	92	RTA00000412F.l.04.1	M00003989D:F12	66372
848	2/24/98	93	RTA00000413F.p.24.1	M00005139A:H03	0
849	2/24/98	94	RTA00000406F.a.23.1	M00003867B:D10	38712
850	2/24/98	95	RTA00000423F.h.05.1	M00003906A:F04	14837
851	2/24/98	96	RTA00000120A.n.19.3	M00001467A:H07	80004
852	2/24/98	97	RTA00000403F.e.01.1	M00001473A:C11	38965
853	2/24/98	98	RTA00000411F.1.03.1	M00003854D:A12	62702
854	2/24/98	99	RTA00000420F.m.19.1	M00005254D:B08	0
855	2/24/98	100	RTA00000339F.o.23.1	M00001473C:D09	7801
856	2/24/98	101	RTA00000121A.m.2.1	M00001507A:A11	81064
857	2/24/98	102	RTA00000420F.g.06.1	M00004891C:D04	0
858	2/24/98	103	RTA00000418F.j.12.1	M00001626C:G08	73316
859	2/24/98	104	RTA00000421F.n.03.1	M00001675C:A04	1638
860	2/24/98	105	RTA00000346F.d.08.1	M00001671A:A10	39955
861	2/24/98	106	RTA00000339F.f.11.1	M00001391C:H02	5832
862	2/24/98	107	RTA00000125A.g.16.1	M00001544A:C09	21497
863	2/24/98	108	RTA00000418F.o.18.1	M00001661B:F06	78676
864	2/24/98	109	RTA00000422F.p.24.2	M00001658A:G09	5823
865	2/24/98	110	RTA00000408F.k.14.1	M00001486B:E12	73856
866	2/24/98	111	RTA00000128A.i.20.1	M00001560A:F03	9900
867	2/24/98	112	RTA00000422F.c.11.1	M00003841D:A04	2643
868	2/24/98	113	RTA00000401F.e.02.1	M00003805B:C04	0
869	2/24/98	114	RTA00000341F.m.21.1	M00004051D:E01	0
870	2/24/98	115	RTA00000418F.h.19.1	M00001590B:C05	0
871	2/24/98	116	RTA00000403F.o.15.1	M00001582B:E12	39140
872	2/24/98	117	RTA00000341F.m.13.1	M00003987B:E12	26502
873	2/24/98	118	RTA00000408F.h.03.1	M00001479D:H03	78382
874	2/24/98	119	RTA00000423F.k.05.1	M00004036D:F02	37472
875	2/24/98	120	RTA00000401F.m.02.1	M00003907A:F01	1573
876	2/24/98	121	RTA00000418F.p.19.1	M00001677D:B01	78544
877	2/24/98	122	RTA00000420F.f.06.1	M00004115D:D08	64812
878	2/24/98	123	RTA00000122A.j.18.1	M00001516A:D05	81317
879	2/24/98	124	RTA00000420F.d.05.1	M00004092B:E05	64432
880	2/24/98	125	RTA00000403F.m.18.1	M00001576A:B09	39185
881	2/24/98	126	RTA00000422F.j.20.1	M00001653A:G07	22388
882	2/24/98	127	RTA00000411F.j.05.1	M00003841C:F06	40709
883	2/24/98	128	RTA00000403F.a.04.1	M00001448A:B12	23529
884	2/24/98	129	RTA00000118A.d.24.1	M00001416A:H02	81488
885	2/24/98	130	RTA00000406F.f.12.1	M00003879A:C11	21895
886	2/24/98	131	RTA00000418F.g.22.1	M00001585B:F01	74837
887	2/24/98	132	RTA00000418F.m.05.1	M00001650B:C10	73600
888	2/24/98	133	RTA00000404F.I.20.1	M00001639B:H05	38638
889	2/24/98	134	RTA00000408F.i.08.2	M00001482A:H05	75811

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
890	2/24/98	135	RTA00000122A.d.5.1	M00001513A:F05	81155
891	2/24/98	136	RTA00000419F.l.12.1	M00003901C:B01	75710
892	2/24/98	137	RTA00000339R.a.06.1	M00001346A:E04	58694
893	2/24/98	138	RTA00000406F.f.03.1	M00003878C:D08	38687
894	2/24/98	139	RTA00000419F.b.19.1	M00003809A:C01	65534
895	2/24/98	140	RTA00000128A.j.6.2	M00001560A:H10	5316
896	2/24/98	141	RTA00000418F.k.19.1	M00001639C:C02	74932
897	2/24/98	142	RTA00000420F.j.19.1	M00005140C:B10	0
898	2/24/98	143	RTA00000420F.h.13.1	M00004899D:G06	0
899	2/24/98	144	RTA00000349R.f.15.1	M00001472A:D08	75097
900	2/24/98	145	RTA00000419F.g.12.1	M00003842C:G03	66171
901	2/24/98	146	RTA00000404F.n.11.2	M00001649A:E11	38001
902	2/24/98	147	RTA00000422F.c.02.1	M00004118B:A03	2902
903	2/24/98	148	RTA00000419F.n.04.1	M00003975C:F07	13102
904	2/24/98	149	RTA00000419F.o.24.1	M00004031A:F07	65092
905	2/24/98	150	RTA00000419F.k.19.1	M00003877C:G12	75447
906	2/24/98	151	RTA00000341F.c.21.1	M00003789C:F06	7899
907	2/24/98	152	RTA00000127A.i.20.1	M00001555A:B12	81418
908	2/24/98	153	RTA00000422F.g.22.1	M00001585B:A06	22561
909	2/24/98	154	RTA00000340F.b.21.1	M00001533D:A08	8001
910	2/24/98	155	RTA00000413F.h.13.1	M00004107A:D01	65190
911	2/24/98	156	RTA00000125A.k.1.1	M00001545A:B12	0
912	2/24/98	157	RTA00000339F.a.23.1	M00001361B:C07	4022
913	2/24/98	158	RTA00000348R.j.16.1	M00001410A:D07	7005
914	2/24/98	159	RTA00000348R.j.17.1	M00001391D:C06	2641
915	2/24/98	160	RTA00000414F.f.19.1	M00005260B:E11	0
916	2/24/98	161	RTA00000418F.n.22.1	M00001659D:B05	79062
917	2/24/98	162	RTA00000406F.l.08.1	M00003908D:D12	39016
918	2/24/98	163	RTA00000422F.l.23.1	M00001616D:C11	4240
919	2/24/98	164	RTA00000345F.k.06.1	M00001475A:A12	0
920	2/24/98	165	RTA00000409F.j.07.1	M00001611C:H11	75190
921	2/24/98	166	RTA00000418F.m.19.1	M00001654D:A03	8890
922	2/24/98	167	RTA00000399F.l.14.1	M00001590B:G08	3354
923	2/24/98	168	RTA00000411F.e.22.1	M00003812B:D07	63638
924	2/24/98	169	RTA00000347F.a.17.1	M00001366D:C06	16723
925	2/24/98	170	RTA00000422F.n.08.1	M00001632B:E05	38655
926	2/24/98	171	RTA00000404F.n.20.1	M00001650A:C11	26865
927	2/24/98	172	RTA00000420F.i.17.1	M00005101C:B09	0
928	2/24/98	173	RTA00000418F.d.13.1	M00001570A:H01	74309
929	2/24/98	174	RTA00000404F.b.02.1	M00001591B:B12	38984
930	2/24/98	175	RTA00000410F.d.09.1	M00001635B:H01	76964
931	2/24/98	176	RTA00000403F.b.10.1	M00001455C:G07	73268
932	2/24/98	177	RTA00000406F.i.12.1	M00003903D:H11	39080
933	2/24/98	178	RTA00000406F.h.08.1	M00003901C:A08	16228
934	2/24/98	179	RTA00000418F.i.19.1	M00001596D:E03	79180
935	2/24/98	180	RTA00000400F.j.19.1	M00001653C:D10	4086
936	2/24/98	181	RTA00000412F.h.21.1	M00003974D:F02	64348
937	2/24/98	182	RTA00000404F.g.14.1	M00001614D:B08	8858
938	2/24/98	183	RTA00000120A.g.18.1	M00001465A:C12	81255
939	2/24/98	184	RTA00000133A.j.13.1	M00001507A:B02	16846
			152		

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln		1400002002C.C05	27059
940	2/24/98	185	RTA00000423F.j.05.1	M00003903C:C05	37958
941	2/24/98	186	RTA00000132A.k.6.1	M00001464A:E07	81284 3077
942	2/24/98	187	RTA00000351R.g.11.1	M00003779D:E08	3077 37458
943	2/24/98	188	RTA00000406F.p.04.1	M00004030D:F11 M00001402D:F02	22446
944	2/24/98	189	RTA00000347F.a.13.1	*	64748
945	2/24/98	190	RTA00000419F.p.23.1	M00004039B:A05	64353
946	2/24/98	191	RTA00000419F.d.17.1	M00003828B:F09	
947	2/24/98	192	RTA00000421F.k.15.1	M00001613D:B03	2222
948	2/24/98	193	RTA00000347F.b.10.1	M00001546C:C07	8044
949	2/24/98	194	RTA00000124A.k.5.1	M00001538A:F12	80252
950	2/24/98	195	RTA00000404F.h.22.1	M00001619C:C07	18735
951	2/24/98	196	RTA00000418F.k.10.1	M00001639A:G07	74454
952	2/24/98	197	RTA00000410F.o.05.1	M00001669A:B02	75262
953	2/24/98	198	RTA00000339R.I.14.1	M00001452A:C07	19119
954	2/24/98	199	RTA00000403F.m.13.2	M00001575D:A10	39077
955	2/24/98	200	RTA00000339F.c.02.1	M00001381C:B08	12975
956	2/24/98	201	RTA00000404F.1.09.1	M00001638B:E12	39176
957	2/24/98	202	RTA00000419F.g.22.1	M00003845D:A09	64515
958	2/24/98	203	RTA00000404F.g.21.1	M00001615C:A11	37947
959	2/24/98	204	RTA00000351R.k.19.1	M00003841B:E03	936
960	2/24/98	205	RTA00000138A.n.4.1	M00001624A:G11	21920
961	2/24/98	206	RTA00000410F.b.15.1	M00001633C:F09	77100
962	2/24/98	207	RTA00000414F.b.08.1	M00005212C:H02	0
963	2/24/98	208	RTA00000419F.j.23.1	M00003871A:C11	74470
964	2/24/98	209	RTA00000411F.j.02.1	M00003841C:D07	65310
965	2/24/98	210	RTA00000419F.p.24.1	M00004039B:E12	63477
966	2/24/98	211	RTA00000404F.a.19.1	M00001590B:C07	38624
967	2/24/98	212	RTA00000408F.k.06.1	M00001485C:H10	78393
968	2/24/98	213	RTA00000123A.f.3.1	M00001531A:H07	44017
969	2/24/98	214	RTA00000404F.h.19.1	M00001619A:E05	8096
970	2/24/98	215	RTA00000403F.j.18.1	M00001539D:E10	5790
971	2/24/98	216	RTA00000420F.i.18.1	M00005101C:E09	0
972	2/24/98	217	RTA00000399F.o.17.1	M00001599D:A09	1106
973	2/24/98	218	RTA00000346F.e.13.1	M00001660B:D03	74653
974	2/24/98	219	RTA00000419F.c.18.1	M00003819D:B11	41394
975	2/24/98	220	RTA00000413F.k.02.1	M00004690A:G08	0
976	2/24/98	221	RTA00000414F.f.13.1	M00005259D:H08	0
977	2/24/98	222	RTA00000405F.e.09.1	M00001663C:F12	38978
978	2/24/98	223	RTA00000404F.e.22.1	M00001610A:H05	11344
979	2/24/98	224	RTA00000341F.g.21.1	M00003914C:F09	8823
980	2/24/98	225	RTA00000414F.d.07.1	M00005229D:H09	0
981	2/24/98	226	RTA00000125A.k.10.1	M00001545A:F02	81644
982	2/24/98	227	RTA00000347F.c.06.1	M00001444D:C01	18846
983	2/24/98	228	RTA00000411F.k.19.1	M00003852D:E08	64200
984	2/24/98	229	RTA00000345F.i.09.1	M00001450A:D08	27250
985	2/24/98	230	RTA00000423F.k.01.1	M00004034D:E09	40426
986	2/24/98	231	RTA00000408F.d.06.1	M00001458D:C11	78997
987	2/24/98	232	RTA00000128A.b.20.1	M00001558A:G09	79761
988	2/24/98	233	RTA00000403F.i.08.1	M00001485C:B10	6176
989	2/24/98	234	RTA00000195AF.d.4.1	M00003881D:D06	22766

SEQ ID NO:	Filing Date of Priority Appln	SEQ ID NO: in Priority Appln	Sequence Name	Clone Name	Cluster ID
989	1/28/98	185	RTA00000195AF.d.4.1	M00003881D:D06	22766
990	2/24/98	235	RTA00000126A.o.23.1	M00001551A:B10	6268
991	2/24/98	236	RTA00000403F.h.12.1	M00001483C:G09	15205
992	2/24/98	237	RTA00000119A.j.22.1	M00001460A:F07	80336
993	2/24/98	238	RTA00000340F.j.12.1	M00001624A:B06	3277
994	2/24/98	239	RTA00000346F.j.02.1	M00003832B:E01	5294
995	2/24/98	240	RTA00000126A.n.7.2	M00001551A:D06	79557
996	2/24/98	241	RTA00000339F.d.13.1	M00001395C:F11	0
997	2/24/98	242	RTA00000404F.j.08.1	M00001629B:B08	39066
998	2/24/98	243	RTA00000410F.c.14.1	M00001634A:H05	77809
999	2/24/98	244	RTA00000120A.g.23.1	M00001465A:E10	81189
1000	2/24/98	245	RTA00000195AF.d.20.1	M00004117A:D11	37574
1000	1/28/98	87	RTA00000195AF.d.20.1	M00004117A:D11	37574
1001	2/24/98	246	RTA00000414F.c.14.1	M00005218A:G05	0
1002	2/24/98	247	RTA00000412F.j.17.1	M00003982C:G04	64071
1003	2/24/98	248	RTA00000404F.k.24.1	M00001636A:C03	15256
1004	2/24/98	249	RTA00000119A.j.10.1	M00001460A:C10	79646
1005	2/24/98	250	RTA00000410F.o.12.1	M00001669A:G12	77376
1006	2/24/98	251	RTA00000119A.i.9.1	M00001457A:G03	0
1007	2/24/98	252	RTA00000412F.g.24.1	M00003973C:C03	28741
1008	2/24/98	253	RTA00000400F.f.18.1	M00001637A:E10	3764
1009	2/24/98	254	RTA00000341F.I.15.1	M00003986B:A08	5294
1010	2/24/98	255	RTA00000419F.o.16.1	M00003989C:G05	62867
1011	2/24/98	256	RTA00000404F.m.03.2	M00001640A:H02	11799
1012	2/24/98	257	RTA00000411F.c.17.1	M00001678D:G03	77664
1013	2/24/98	258	RTA00000406F.k.15.1	M00003907C:C04	38549
1014	2/24/98	259	RTA00000406F.a.02.1	M00003855C:F10	37744
1015	2/24/98	260	RTA00000414F.e.08.1	M00005236A:E04	0
1016	2/24/98	261	RTA00000341F.b.06.1	M00003794A:E12	17008
1017	2/24/98	262	RTA00000409F.n.14.1	M00001621B:G05	78190
1018	2/24/98	263	RTA00000410F.p.17.1	M00001674D:C10	47425
1019	2/24/98	264	RTA00000345F.j.08.1	M00001451B:A04	16731
1020	2/24/98	265	RTA00000340F.k.16.1	M00001647B:C09	13157
1021	2/24/98	266	RTA00000419F.g.15.1	M00003844D:A07	32519
1022	2/24/98	267	RTA00000423F.a.19.1	M00001676D:A02	21396
1023	2/24/98	268	RTA00000403F.e.23.1	M00001476A:D11	9626
1024	2/24/98	269	RTA00000422F.e.08.1	M00001573A:E01	39020
1025	2/24/98	270	RTA00000411F.d.15.1	M00001692A:B06	74890
1026	2/24/98	271	RTA00000414F.e.16.1	M00005236B:H10	0
1027	2/24/98	272	RTA00000411F.I.15.1	M00003857C:F11	66704
1028	2/24/98	273	RTA00000400F.a.11.1	M00001612B:D11	0
1029	2/24/98	274	RTA00000405F.e.08.1	M00001663C:F10	37916
1030	2/24/98	275	RTA00000353R.j.24.1	M00001428B:D01	23089
1031	2/24/98	276	RTA00000423F.a.18.1	M00001675A:G10	26761
1032	2/24/98	277	RTA00000418F.o.06.1	M00001660C:D11	75930
1033	2/24/98	278	RTA00000404F.c.10.1	M00001593B:E11	23534
1034	2/24/98	279	RTA00000418F.i.21.1	M00001596D:E10	78728
1035	2/24/98	280	RTA00000411F.1.13.1	M00001671C:C11	31066
1036 1037	2/24/98 2/24/98	281 282	RTA00000411F.I.13.1	M00003857C:C09	43114
105/	2124170	404	RTA00000407F.a.24.1	M00004083A:E08	37560

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appin	DT 1 0000001 (F) 0 (1	N400004120C A 12	10420
1038	2/24/98	283	RTA00000346F.n.06.1	M00004139C:A12	12439
1039	2/24/98	284	RTA00000412F.1.21.1	M00004029C:G10	65183
1040	2/24/98	285	RTA00000413F.i.02.1	M00004110D:A10	65857
1041	2/24/98	286	RTA00000404F.i.19.1	M00001625B:C10	38698
1042	2/24/98	287	RTA00000410F.n.09.1	M00001662C:A04	11736
1043	2/24/98	288	RTA00000403F.a.11.1	M00001448C:F10	73109
1044	2/24/98	289	RTA00000420F.n.08.1	M00005257A:H11	0
1045	2/24/98	290	RTA00000411F.k.16.1	M00003852C:B06	64759
1046	2/24/98	291	RTA00000405F.c.01.1	M00001657D:A04	19236
1047	2/24/98	292	RTA00000423F.i.18.1	M00003918A:D08	14996
1048	2/24/98	293	RTA00000403F.l.04.1	M00001571C:A04	39278
1049	2/24/98	294	RTA00000405F.l.17.1	M00003805A:F02	17225
1050	2/24/98	295	RTA00000406F.a.07.1	M00003856C:H09	26607
1051	2/24/98	296	RTA00000347F.d.06.1	M00001457C:F02	39122
1052	2/24/98	297	RTA00000419F.b.18.1	M00003808D:D08	67034
1053	2/24/98	298	RTA00000406F.h.07.1	M00003901B:H04	38003
1054	2/24/98	299	RTA00000405F.I.15.1	M00001694A:E03	19575
1055	2/24/98	300	RTA00000406F.g.17.1	M00003881B:F10	37979
1056	2/24/98	301	RTA00000401F.m.23.1	M00003914C:C02	2801
1057	2/24/98	302	RTA00000356R.f.18.1	M00004692A:H10	0
1058	2/24/98	303	RTA00000130A.h.22.1	M00001617A:D06	80933
1059	2/24/98	304	RTA00000403F.n.18.2	M00001577D:H06	8811
1060	2/24/98	305	RTA00000418F.p.06.1	M00001664A:F08	32628
1061	2/24/98	306	RTA00000404F.d.13.1	M00001595D:A04	39036
1062	2/24/98	307	RTA00000420F.1.12.2	M00005230B:H09	0
1063	2/24/98	308	RTA00000353R.d.11.1	M00004692A:H08	0
1064	2/24/98	309	RTA00000340F.n.01.1	M00001679A:G06	39081
1065	2/24/98	310	RTA00000419F.d.06.1	M00003820B:D07	65496
1066	2/24/98	311	RTA00000419F.n.09.1	M00003977C:A06	66070
1067	2/24/98	312	RTA00000399F.i.08.1	M00001575D:B10	38927
1068	2/24/98	313	RTA00000406F.g.07.1	M00003880C:E11	37925
1069	2/24/98	314	RTA00000423F.g.13.1	M00003905A:E07	38028
1070	2/24/98	315	RTA00000419F.p.12.1	M00004037A:E04	13767
1071	2/24/98	316	RTA00000414F.a.02.1	M00005178D:H04	0
1072	2/24/98	317	RTA00000195AF.b.21.1	M00001595B:A09	39055
1072	1/28/98	602	RTA00000195AF.b.21.1	M00001595B:A09	39055
1073	2/24/98	318	RTA00000403F.h.05.1	M00001482D:A04	39096
1074	2/24/98	319	RTA00000420F.b.21.1	M00004088D:B10	65057
1075	2/24/98	320	RTA00000422F.p.07.2	M00001661A:E06	39024
1076	2/24/98	321	RTA00000339F.c.21.1	M00001389C:A08	5325
1077	2/24/98	322	RTA00000339F.c.24.1	M00001364B:B06	5516
1078	2/24/98	323	RTA00000421F.n.19.1	M00001679A:D10	16409
1079	2/24/98	324	RTA00000340F.p.17.1	M00003750C:H05	0
1080	2/24/98	325	RTA00000345F.k.21.1	M00001464B:C11	40204
1081	2/24/98	326	RTA00000419F.b.15.1	M00003806D:D11	43969
1082	2/24/98	327	RTA00000405F.a.11.1	M00001655A:B11	39124
1083	2/24/98	328	RTA00000423F.k.19.2	M00003985D:E10	17615
1084	2/24/98	329	RTA00000413F.e.16.1	M00004093C:C02	63836
1085	2/24/98	330	RTA00000403F.i.04.1	M00001485B:D09	8930
1086	2/24/98	331	RTA00000404F.o.18.2	M00001651C:C05	39110

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
1087	2/24/98	332	RTA00000409F.i.24.1	M00001611B:A09	76967
1088	2/24/98	333	RTA00000399F.f.11.1	M00001487C:F01	40167
1089	2/24/98	334	RTA00000408F.p.05.1	M00001575B:B02	9649
1090	2/24/98	335	RTA00000413F.d.02.1	M00004087B:A12	66172
1091	2/24/98	336	RTA00000340F.n.13.1	M00001688D:B10	17055
1092	2/24/98	337	RTA00000340F.p.04.1	M00001679D:B02	78533
1093	2/24/98	338	RTA00000411F.c.05.1	M00001677B:H06	73368
1094	2/24/98	339	RTA00000403F.g.10.1	M00001481A:G06	20211
1095	2/24/98	340	RTA00000408F.I.13.1	M00001530A:B12	4423
1096	2/24/98	341	RTA00000412F.g.20.2	M00003972C:F08	25018
1097	2/24/98	342	RTA00000404F.i.02.1	M00001619D:D10	39015
1098	2/24/98	343	RTA00000422F.g.21.1	M00001583A:F07	17232
1099	2/24/98	344	RTA00000403F.m.15.2	M00001575D:D12	26901
1100	2/24/98	345	RTA00000412F.h.23.2	M00003974D:H04	65118
1101	2/24/98	346	RTA00000418F.j.08.1	M00001626C:C11	73382
1102	2/24/98	347	RTA00000125A.n.4.1	M00001546A:D08	81984
1103	2/24/98	348	RTA00000412F.l.19.1	M00004029C:C05	65825
1104	2/24/98	349	RTA00000404F.m.10.2	M00001641D:E02	779
1105	2/24/98	350	RTA00000129A.p.3.1	M00001604A:B08	32644
1106	2/24/98	351	RTA00000340F.p.20.1	M00003752B:C02	17008
1107	2/24/98	352	RTA00000411F.a.10.1	M00001675C:G01	73073
1108	2/24/98	353	RTA00000409F.n.17.1	M00001621C:C10	76725
1109	2/24/98	354	RTA00000404F.c.03.2	M00001592C:F11	39198
1110	2/24/98	355	RTA00000420F.a.19.1	M00004076A:D12	34192
1111	2/24/98	356	RTA00000409F.m.24.1	M00001620D:H02	3942
1112	2/24/98	357	RTA00000406F.n.16.1	M00003972A:G09	5660
1113	2/24/98	358	RTA00000414F.e.06.1	M00005235A:A03	0
1114	2/24/98	359	RTA00000420F.d.12.1	M00004096D:H03	64095
1115	2/24/98	360	RTA00000409F.j.19.1	M00001613A:F03	73792
1116	2/24/98	361	RTA00000422F.d.16.1	M00001570C:G03	39133
1117	2/24/98	362	RTA00000418F.m.16.1	M00001653B:E06	74986
1118	2/24/98	363	RTA00000405F.c.11.1	M00001659A:D12	39068
1119	2/24/98	364	RTA00000404F.k.22.1	M00001635D:C12	39084
1120	2/24/98	365	RTA00000418F.k.07.1	M00001637A:F10	75067
1121	2/24/98	366	RTA00000403F.c.10.1	M00001456D:F05	75261
1122	2/24/98	367	RTA00000401F.o.06.1	M00004029C:C12	2679
1123	2/24/98	368	RTA00000346F.o.08.1	M00004149C:B02	0
1124	2/24/98	369	RTA00000410F.m.05.1	M00001657B:B04	74964
1125	2/24/98	370	RTA00000405F.i.20.1	M00001677A:G11	38532
1126	2/24/98	371	RTA00000403F.j.17.1	M00001539D:B10	38563
1127	2/24/98	372	RTA00000408F.p.24.1	M00001579A:E03	74286
1128	2/24/98	373	RTA00000418F.k.18.1	M00001639C:A10	75385
1129	2/24/98	374	RTA00000422F.m.04.1	M00001615B:A09	38702
1130	2/24/98	375	RTA00000405F.g.16.2	M00001672D:D04	9021
1131	2/24/98	376	RTA00000400F.k.22.1	M00001656A:B07	2512
1132	2/24/98	377	RTA00000346F.i.01.1	M00003797A:D06	22260
1133	2/24/98	378	RTA00000403F.a.07.1	M00001448B:F09	73559
1134	2/24/98	379	RTA00000349R.j.07.1	M00001529B:C04	2642
1135	2/24/98	380	RTA00000403F.b.19.1	M00001456B:A06	22327
1136	2/24/98	381	RTA00000418F.m.23.1	M00001654D:F11	77195

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
1137	2/24/98	382	RTA00000341F.h.10.1	M00003901B:G11	0
1138	2/24/98	383	RTA00000404F.i.18.1	M00001621C:H12	21912
1139	2/24/98	384	RTA00000422F.i.14.1	M00001487A:F10	39300
1140	2/24/98	385	RTA00000418F.m.14.1	M00001651B:E06	75711
1141	2/24/98	386	RTA00000406F.o.12.1	M00003986D:D02	37459
1142	2/24/98	387	RTA00000411F.a.15.1	M00001675D:B08	73812
1143	2/24/98	388	RTA00000411F.a.07.1	M00001675C:C03	74547
1144	2/24/98	389	RTA00000411F.c.02.1	M00001677B:B04	72852
1145	2/24/98	390	RTA00000355R.a.14.1	M00004187D:G09	10207
1146	2/24/98	391	RTA00000130A.h.16.1	M00001617A:A08	80761
1147	2/24/98	392	RTA00000410F.p.23.1	M00001675B:C01	73948
1148	2/24/98	393	RTA00000418F.m.24.1	M00001654D:F12	77114
1149	2/24/98	394	RTA00000420F.m.02.1	M00005233A:G08	0
1150	2/24/98	395	RTA00000408F.j.19.2	M00001485C:C08	73752
1151	2/24/98	396	RTA00000406F.e.21.1	M00003877D:G05	9090
1152	2/24/98	397	RTA00000118A.d.17.1	M00001416A:D09	81921
1153	2/24/98	398	RTA00000407F.b.04.1	M00004086D:B09	63221
1154	2/24/98	399	RTA00000411F.e.07.1	M00003810C:A03	65008
1155	2/24/98	400	RTA00000403F.f.08.1	M00001477A:G07	19107
1156	2/24/98	401	RTA00000132A.c.11.1	M00001454A:G03	87278
1157	2/24/98	402	RTA00000420F.e.16.1	M00004110A:E04	63639
1158	2/24/98	403	RTA00000403F.d.22.1	M00001473A:A07	10692
1159	2/24/98	404	RTA00000404F.b.11.1	M00001591D:F06	39079
1160	2/24/98	405	RTA00000418F.k.17.1	M00001639C:A09	75390
1161	2/24/98	406	RTA00000129A.k.12.1	M00001601A:A06	79322
1162	2/24/98	407	RTA00000340R.m.07.1	M00001679D:F02	78415
1163	2/24/98	408	RTA00000405F.d.14.1	M00001662A:C12	35209
1164	2/24/98	409	RTA00000406F.f.11.1	M00003879A:B08	38601
1165	2/24/98	410	RTA00000120A.h.5.1	M00001465A:G06	80344
1166	2/24/98	411	RTA00000420F.m.12.1	M00005234D:B04	0
1167	2/24/98	412	RTA00000411F.g.06.1	M00003822D:C06	66065
1168	2/24/98	413	RTA00000408F.d.16.1	M00001459B:D03	76318
1169	2/24/98	414	RTA00000120A.p.18.1	M00001468A:C05	6478
1170	2/24/98	415	RTA00000340R.f.05.1	M00001569B:G11	3202
1171	2/24/98	416	RTA00000404F.c.19.1	M00001594A:D06	39026
1172	2/24/98	417	RTA00000423F.1.02.1	M00003978C:A03	5639
1173	2/24/98	418	RTA00000410F.a.01.1	M00001631D:B10	73354
1174	2/24/98	419	RTA00000408F.h.08.1	M00001480A:D03	74575
1175	2/24/98	420	RTA00000422F.b.16.1	M00003813B:A11	17045
1176	2/24/98	421	RTA00000419F.f.10.1	M00003835D:G06	66193
1177	2/24/98	422	RTA00000418F.l.04.1	M00001641C:D02	74140
1178	2/24/98	423	RTA00000410F.a.16.1	M00001633A:E06	73548
1179	2/24/98	424	RTA00000138A.e.13.1	M00001605A:E06	79608
1180	2/24/98	425	RTA00000130A.b.5.1	M00001605A:E09	79579
1181	2/24/98	426	RTA00000408F.j.15.2	M00001485B:F05	74759
1182	2/24/98	427	RTA00000410F.m.20.1	M00001660B:E03	74285
1183	2/24/98	428	RTA00000422F.f.14.1	M00001478B:D07	2036
1184	2/24/98	429	RTA00000422F.c.17.1	M00004099D:F01	1360
1185	2/24/98	430	RTA00000419F.e.04.1	M00003831C:G05	62963
1186	2/24/98	431	RTA00000399F.j.15.1	M00001578C:G06	1261
			157		

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
1187	2/24/98	432	RTA00000418F.g.05.1	M00001579C:H06	73075
1188	2/24/98	433	RTA00000419F.n.02.1	M00003958B:H08	65963
1189	2/24/98	434	RTA00000348R.b.16.1	M00001347B:H04	6510
1190	2/24/98	435	RTA00000340F.b.02.1	M00001503C:G05	10185
1191	2/24/98	436	RTA00000119A.m.15.1	M00001461A:E05	80989
1192	2/24/98	437	RTA00000403F.m.20.2	M00001576A:F11	707
1193	2/24/98	438	RTA00000195R.d.09.1	M00003981C:B04	8537
1194	2/24/98	439	RTA00000413F.g.23.1	M00004103B:E09	40700
1195	2/24/98	440	RTA00000403F.a.18.1	M00001448D:F12	75726
1196	2/24/98	441	RTA00000404F.m.20.2	M00001647A:H08	39144
1197	2/24/98	442	RTA00000347F.b.02.1	M00001450A:A02	39304
1198	2/24/98	443	RTA00000414F.f.15.1	M00005260A:A12	0
1199	2/24/98	444	RTA00000419F.h.04.1	M00003846A:D03	65034
1200	2/24/98	445	RTA00000408F.d.12.1	M00001459B:A12	75782
1201	2/24/98	446	RTA00000133A.m.19.2	M00001512A:G05	80167
1202	2/24/98	447	RTA00000423F.b.04.3	M00001675D:E10	6311
1203	2/24/98	448	RTA00000127A.a.3.1	M00001552A:H10	13232
1203	2/24/98	449	RTA00000411F.j.16.1	M00003843A:E08	17237
1205	2/24/98	450	RTA00000118A.a.23.1	M00001395A:H02	3500
1206	2/24/98	451	RTA00000126A.o.22.1	M00001551A:A11	81752
1207	2/24/98	452	RTA00000419F.n.13.1	M00003977D:A06	66026
1208	2/24/98	453	RTA00000130A.h.13.1	M00001617A:A01	80790
1209	2/24/98	454	RTA00000418F.n.19.1	M00001659C:F02	28761
1210	2/24/98	455	RTA00000399F.d.23.1	M00001481B:A07	3310
1211	2/24/98	456	RTA00000413F.o.06.1	M00005100A:B02	0
1212	2/24/98	457	RTA00000411F.m.19.1	M00003868D:D11	74924
1213	2/24/98	458	RTA00000130A.a.19.1	M00001605A:A06	0
1214	2/24/98	459	RTA00000419F.k.06.1	M00003871D:A10	78493
1215	2/24/98	460	RTA00000341F.j.12.1	M00003987C:G03	12195
1216	2/24/98	461	RTA00000412F.d.16.1	M00003906B:H06	26829
1217	2/24/98	462	RTA00000119A.j.23.1	M00001460A:G07	79835
1218	2/24/98	463	RTA00000403F.o.22.1	M00001583A:D01	25076
1219	2/24/98	464	RTA00000195AF.c.12.1	M00003818B:G12	37582
1219	1/28/98	300	RTA00000195AF.c.12.1	M00003818B:G12	37582
1220	2/24/98	465	RTA00000350R.p.18.1	M00001676B:F05	11460
1221	2/24/98	466	RTA00000406F.i.24.1	M00003904D:B12	12767
1222	2/24/98	467	RTA00000123A.n.13.2	M00001534A:D03	39167
1223	2/24/98	468	RTA00000423F.c.19.1	M00001680B:E10	40472
1224	2/24/98	469	RTA00000405F.g.24.1	M00001673D:D06	39076
1225	2/24/98	470	RTA00000411F.j.06.1	M00003841C:H08	63545
1226	2/24/98	471	RTA00000419F.c.11.1	M00003817B:C04	65504
1227	2/24/98	472	RTA00000135A.f.14.2	M00001542A:G12	79969
1228	2/24/98	473	RTA00000403F.a.05.1	M00001448A:E11	18808
1229	2/24/98	474	RTA00000405F.e.17.1	M00001669A:C10	38662
1230	2/24/98	475	RTA00000411F.d.05.1	M00001681C:A08	75812
1231	2/24/98	476	RTA00000345F.h.01.1	M00001441B:D11	10834
1232	2/24/98	477	RTA00000418F.d.03.1	M00001567B:G11	76824
1233	2/24/98	478	RTA00000418F.h.08.1	M00001589B:E07	76401
1234	2/24/98	479	RTA00000418F.m.10.1	M00001651A:H11	79110
1235	2/24/98	480	RTA00000411F.i.15.1	M00003837C:G08	31612

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
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1236	2/24/98	481	RTA00000413F.i.23.1	M00004118B:F01	63073
1237	2/24/98	482	RTA00000411F.e.24.1	M00003813A:B02	64781
1238	2/24/98	483	RTA00000406F.g.22.1	M00003881D:C12	38590
1239	2/24/98	484	RTA00000126A.n.13.2	M000035612:012	79735
1240	2/24/98	485	RTA00000419F.a.02.1	M00001678A:F05	77993
1240	2/24/98	486	RTA00000346F.l.13.1	M00003980B:C11	7542
1241	2/24/98	487	RTA00000340F.g.05.1	M00003300B:C11	0
1242	2/24/98	488	RTA000004201.g.03.1 RTA00000339F.k.23.1	M00001429D:H12	Ö
1243	2/24/98	489	RTA000003551.ki.25.1	M00003906A:F12	1685
1244	2/24/98	490	RTA00000120A.d.15.1	M0000336071172	80533
1245	2/24/98	491	RTA00000418F.f.21.1	M00001579B:F04	75157
1247	2/24/98	492	RTA00000340F.o.18.1	M00001579D:101	4261
1247	2/24/98	493	RTA000003401.5.16.1 RTA00000129A.d.1.2	M00001587A:F05	80058
1248	2/24/98	494	RTA0000012971.d.11.2	M0000130771:103 M00003876C:F02	0
1249	2/24/98	495	RTA00000419F.R.12.11	M00003676C:F02	16259
1250	2/24/98	496	RTA000004001.0.21.11 RTA00000419F.m.20.1	M000010076:200 M00003914A:B07	76720
1251	2/24/98	497	RTA000004771.iii.20.1	M0000351471:B07	22110
1252	2/24/98	498	RTA00000330R.1.21.1 RTA00000406F.e.15.1	M00001010C:207	39074
1255	2/24/98	499	RTA000004001.c.13.11 RTA00000126A.p.18.2	M00003677C:X11	80881
1254	2/24/98	500	RTA00000120A.p.10.2	M00001532A:E10	73117
1256	2/24/98	501	RTA000004111.c.10.1	M00001078D:B11	0
1257	2/24/98	502	RTA000004141.1.03.1 RTA00000341F.d.08.1	M00003237D:1111 M00003824C:D07	0
1257	2/24/98	503	RTA000003411.d.08.1	M00005233B:D04	0
1259	2/24/98	504	RTA000004201.iii.00.1	M00003233B:D04	64788
1260	2/24/98	505	RTA000004131.d.03.1	M00004007C:707	81437
1261	2/24/98	506	RTA0000012174.0.3.1 RTA00000403F.f.09.1	M000013171:7102 M00001477B:C02	0
1262	2/24/98	507	RTA000004031.1.07.1 RTA00000420F.e.02.1	M00001477B:C02 M00004107B:D07	40259
1263	2/24/98	508	RTA00000420F.i.20.1	M00005101C:E12	0
1264	2/24/98	509	RTA00000349R.g.10.1	M00003101C:E12	5777
1265	2/24/98	510	RTA00000131A.g.16.2	M00001449A:F01	0
1266	2/24/98	511	RTA00000341F.b.13.1	M00003762B:H09	Ŏ
1267	2/24/98	512	RTA00000414F.c.16.1	M00005228A:B03	0
1268	2/24/98	513	RTA00000126A.k.7.2	M00001550A:E07	79866
1269	2/24/98	514	RTA00000404F.e.13.1	M00001608D:E09	12046
1270	2/24/98	515	RTA00000419F.I.03.1	M00003879A:D02	79060
1271	2/24/98	516	RTA00000339F.f.20.1	M00001399A:C03	6494
1272	2/24/98	517	RTA00000118A.a.2.1	M00001395A:A12	38067
1273	2/24/98	518	RTA00000410F.m.18.1	M00001660B:A09	76365
1274	2/24/98	519	RTA00000404F.1.10.1	M00001638B:F10	23136
1275	2/24/98	520	RTA00000406F.c.20.1	M00003871D:G06	38578
1276	2/24/98	521	RTA00000413F.b.14.1	M00004078A:C11	66591
1277	2/24/98	522	RTA00000406F.c.18.1	M00003871C:F12	14368
1278	2/24/98	523	RTA00000418F.j.09.1	M00001626C:D12	76352
1279	2/24/98	524	RTA00000419F.f.23.1	M00003840D:H10	65002
1280	2/24/98	525	RTA00000348R.d.24.1	M00001349B:G05	5774
1281	2/24/98	526	RTA00000411F.a.05.1	M00001675B:H03	76699
1282	2/24/98	527	RTA00000419F.m.21.1	M00003914A:E04	77947
1283	2/24/98	528	RTA00000405F.n.16.1	M00003825B:B10	21503
1284	2/24/98	529	RTA00000422F.o.19.2	M00001655C:E01	13084
1285	2/24/98	530	RTA00000408F.n.02.2	M00001539A:E01	76993

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
1206	Appln 2/24/98	Appln 531	RTA00000345F.n.12.1	M00001528A:C04	7337
1286		532	RTA00000343F.ii.12.1 RTA00000403F.a.24.1	M00001328A.C04 M00001455B:A09	24128
1287	2/24/98		RTA00000403F.a.24.1 RTA00000423F.e.11.1	M00001453B.A09 M00003809B:E10	2566
1288	2/24/98	533		M00003809B.E10 M00001548A:H04	1902
1289	2/24/98	534	RTA00000110A = 7.1	M00001348A:F11	83580
1290	2/24/98	535	RTA00000119A.g.7.1	M00001434A.F11 M00003835B:H11	66975
1291	2/24/98	536	RTA00000411F.i.02.1		75487
1292	2/24/98	537	RTA00000408F.l.09.1	M00001530A:A09	23012
1293	2/24/98	538	RTA00000423F.g.04.1	M00003903D:C12	
1294	2/24/98	539	RTA00000346F.m.15.1	M00004037B:C04	13553
1295	2/24/98	540	RTA00000418F.i.18.1	M00001595C:B05	78024
1296	2/24/98	541	RTA00000411F.h.15.1	M00003832A:A09	65160
1297	2/24/98	542	RTA00000410F.i.19.1	M00001641B:C10	78988
1298	2/24/98	543	RTA00000419F.k.24.1	M00003878C:G08	75596
1299	2/24/98	544	RTA00000420F.l.21.2	M00005232A:H12	0
1300	2/24/98	545	RTA00000420F.e.15.1	M00004110A:A10	20190
1301	2/24/98	546	RTA00000409F.i.09.1	M00001610B:C07	75279
1302	2/24/98	547	RTA00000419F.h.02.1	M00003845D:G08	63985
1303	2/24/98	548	RTA00000413F.b.12.1	M00004077B:H11	64932
1304	2/24/98	549	RTA00000121A.h.18.1	M00001471A:B04	16376
1305	2/24/98	550	RTA00000411F.n.20.1	M00003875C:A09	75816
1306	2/24/98	551	RTA00000340F.b.05.1	M00001513A:G07	0
1307	2/24/98	552	RTA00000411F.n.12.1	M00003875A:C04	73308
1308	2/24/98	553	RTA00000408F.j.12.2	M00001485B:C03	18226
1309	2/24/98	554	RTA00000409F.i.03.1	M00001610A:E09	75968
1310	2/24/98	555	RTA00000133A.d.22.1	M00001469A:G11	11797
1311	2/24/98	556	RTA00000400F.i.11.1	M00001649C:H10	2587
1312	2/24/98	557	RTA00000409F.j.05.1	M00001611C:C12	74128
1313	2/24/98	558	RTA00000419F.m.04.1	M00003906C:C05	74367
1314	2/24/98	559	RTA00000418F.k.03.1	M00001634D:G11	78901
1315	2/24/98	560	RTA00000419F.d.16.1	M00003828B:E07	64357
1316	2/24/98	561	RTA00000420F.e.10.1	M00004108D:G04	65899
1317	2/24/98	562	RTA00000401F.j.17.1	M00003901B:C05	5483
1318	2/24/98	563	RTA00000406F.b.08.1	M00003867D:A06	18258
1319	2/24/98	564	RTA00000418F.k.08.1	M00001639A:C03	18259
1320	2/24/98	565	RTA00000420F.k.17.2	M00005217B:A06	0
1321	2/24/98	566	RTA00000414F.d.05.1	M00005229D:H03	0
1322	2/24/98	567	RTA00000410F.c.02.1	M00001633D:D12	75055
1323	2/24/98	568	RTA00000403F.m.03.1	M00001573D:D10	39179
1324	2/24/98	569	RTA00000403F.h.18.1	M00001484C:A04	39241
1325	2/24/98	570	RTA00000405F.n.13.1	M00003824A:G10	23810
1326	2/24/98	571	RTA00000355R.e.14.1	M00004314B:G07	16837
1327	2/24/98	572	RTA00000422F.I.03.1	M00001610D:D05	39147
1328	2/24/98	573	RTA00000414F.c.23.1	M00005229B:G12	0
1329	2/24/98	574	RTA00000403F.o.14.1	M00001579D:H09	38971
1330	2/24/98	575	RTA00000345F.a.18.1	M00001351C:B06	5517
1331	2/24/98	576	RTA00000401F.d.15.2	M00001693C:C12	5297
1332	2/24/98	577	RTA00000419F.e.11.1	M00003833B:C12	36780
1333	2/24/98	578	RTA00000127A.f.11.1	M00001554A:A08	81463
1334	2/24/98	579	RTA00000413F.m.16.1	M00004898C:F03	0
1335	2/24/98	580	RTA00000403F.o.07.1	M00001579C:A01	39037

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
1336	2/24/98	581	RTA00000403F.d.19.1	M00001472C:A01	39243
1337	2/24/98	582	RTA00000414F.e.14.1	M00005236B:F10	0
1338	2/24/98	583	RTA00000406F.i.17.1	M00003904B:C03	37902
1339	2/24/98	584	RTA00000418F.d.22.1	M00001573B:C06	75324
1340	2/24/98	585	RTA00000340R.o.12.1	M00003746C:E02	53732
1341	2/24/98	586	RTA00000125A.g.24.1	M00001544A:F05	80397
1342	2/24/98	587	RTA00000130A.o.21.1	M00001623A:F04	80218
1343	2/24/98	588	RTA00000420F.a.23.1	M00004078B:F12	42158
1344	2/24/98	589	RTA00000411F.m.18.1	M00003868D:D09	75629
1345	2/24/98	590	RTA00000407F.b.22.1	M00004108B:B02	37487
1346	2/24/98	591	RTA00000409F.a.16.1	M00001583A:A05	73990
1347	2/24/98	592	RTA00000421F.p.18.1	M00003877B:H10	750
1348	2/24/98	593	RTA00000341F.k.12.1	M00004103C:D04	62985
1349	2/24/98	594	RTA00000129A.c.18.2	M00001587A:B10	37216
1350	2/24/98	595	RTA00000410F.d.10.1	M00001635B:H02	77561
1351	2/24/98	596	RTA00000351R.i.03.1	M00003846B:D06	6874
1351	2/24/98	597	RTA00000135A.l.1.2	M00001545A:B10	39426
1352	2/24/98	598	RTA00000420F.b.18.1	M00004086D:G08	66136
1354	2/24/98	599	RTA00000401F.k.14.1	M00003903A:H09	211
1355	2/24/98	600	RTA00000406F.m.04.1	M00003914B:A11	14959
1356	2/24/98	601	RTA00000403F.o.13.1	M00003571D::F11	39049
1357	2/24/98	602	RTA00000411F.f.06.1	M00001373B:E09	64186
1357	2/24/98	603	RTA00000399F.o.19.1	M00003613B:E03	2594
1359	2/24/98	604	RTA000003511.c.13.1	M00003747D:C05	11476
1360	2/24/98	605	RTA00000331R.c.13.1	M00003717D:203	0
1361	2/24/98	606	RTA00000420F.1.20.2	M00005232A:C10	Ö
1362	2/24/98	607	RTA00000420F.d.16.1	M00004103D:F10	64485
1363	2/24/98	608	RTA00000404F.i.12.1	M000011620D:G11	39001
1364	2/24/98	609	RTA00000404F.o.10.2	M00001651B:B12	16785
1365	2/24/98	610	RTA00000419F.d.07.1	M00003820B:D10	21421
1366	2/24/98	611	RTA00000404F.p.02.2	M00001652D:A06	39097
1367	2/24/98	612	RTA00000125A.k.14.1	M00001545A:G05	79457
1368	2/24/98	613	RTA00000122A.j.22.1	M00001516A:F06	81151
1369	2/24/98	614	RTA00000406F.i.13.1	M00003904A:C04	37904
1370	2/24/98	615	RTA00000135A.b.23.1	M00001538A:D12	35241
1371	2/24/98	616	RTA00000423F.c.11.1	M00001677D:B02	0
1372	2/24/98	617	RTA00000423F.f.23.1	M00003816C:E09	15390
1373	2/24/98	618	RTA00000423F.1.04.1	M00004039B:G08	14320
1374	2/24/98	619	RTA00000420F.b.04.1	M00004081A:E02	63820
1375	2/24/98	620	RTA00000420F.a.07.1	M00004072C:F08	63405
1376	2/24/98	621	RTA00000408F.i.18.2	M00001482C:D02	74410
1377	2/24/98	622	RTA00000404F.1.07.1	M00001637C:C06	10798
1378	2/24/98	623	RTA00000341F.j.05.1	M00003963D:B05	36177
1379	2/24/98	624	RTA00000420F.a.16.1	M00004075D:C10	63345
1380	2/24/98	625	RTA00000126A.h.22.2	M00001549A:F01	0
1381	2/24/98	626	RTA00000410F.j.01.1	M00001641B:F12	73399
1382	2/24/98	627	RTA00000408F.p.21.1	M00001579A:C03	77930
1383	2/24/98	628	RTA00000412F.d.19.1	M00003907B:C03	75743
1384	2/24/98	629	RTA00000352R.c.04.1	M00003924A:D08	71976
1385	2/24/98	630	RTA00000413F.f.19.1	M00004100B:C07	65189

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
1101	Priority	Priority			
	Appln	Appln			
1386	2/24/98	631	RTA00000411F.e.03.1	M00001694D:C12	73648
1387	2/24/98	632	RTA00000191AF.j.14.1	M00004073A:H12	1002
1387	1/28/98	387	RTA00000191AF.j.14.1	M00004073A:H12	1002
1388	2/24/98	633	RTA00000341F.d.02.1	M00003797A:G03	4706
1389	2/24/98	634	RTA00000418F.c.04.1	M00001487B:A11	41587
1390	2/24/98	635	RTA00000418F.o.17.1	M00001661B:F03	79069
1391	2/24/98	636	RTA00000418F.e.21.1	M00001577B:A03	74773
1392	2/24/98	637	RTA00000419F.d.14.1	M00003828A:D05	64945
1393	2/24/98	638	RTA00000418F.b.09.1	M00001478B:H08	19700
1394	2/24/98	639	RTA00000414F.d.09.1	M00005231C:B01	0
1395	2/24/98	640	RTA00000405F.f.02.1	M00001669B:G02	38665
1396	2/24/98	641	RTA00000410F.j.20.1	M00001642D:G10	73601
1397	2/24/98	642	RTA00000341F.h.19.1	M00003916C:C05	0
1398	2/24/98	643	RTA00000420F.1.14.2	M00005230D:F06	0
1399	2/24/98	644	RTA00000119A.j.9.1	M00001460A:B12	82060
1400	2/24/98	645	RTA00000422F.p.12.2	M00001661C:F10	9840
1401	2/24/98	646	RTA00000421F.m.14.1	M00001642A:F03	3524
1402	2/24/98	647	RTA00000418F.b.23.1	M00001485A:C05	28767
1403	2/24/98	648	RTA00000340F.i.13.1	M00001624B:B10	79299
1404	2/24/98	649	RTA00000412F.g.03.1	M00003971B:A10	64740
1405	2/24/98	650	RTA00000122A.g.17.1	M00001514A:B08	32655
1406	2/24/98	651	RTA00000403F.g.11.1	M00001481A:H08	24238
1407	2/24/98	652	RTA00000419F.n.12.1	M00003977D:A03	66086
1408	2/24/98	653	RTA00000352R.m.12.1	M00004212B:C07	2379
1409	2/24/98	654	RTA00000421F.a.05.1	M00001570C:G06	5278
1410	2/24/98	655	RTA00000351R.p.14.1	M00003915C:H04	13166
1411	2/24/98	656	RTA00000403F.e.08.1	M00001473D:B11	19126
1412	2/24/98	657	RTA00000124A.k.20.1	M00001538A:C08	80913
1413	2/24/98	658	RTA00000121A.n.2.1	M00001511A:A05	33585 39159
1414	2/24/98	659	RTA00000422F.m.24.1	M00001641D:C04	75002
1415	2/24/98	660	RTA00000408F.e.24.2	M00001476C:C11	73002 8479
1416	2/24/98	661	RTA00000341F.l.16.1	M00003986D:C08 M00001473D:G01	2566
1417	2/24/98	662	RTA00000339F.o.07.1		78775
1418	2/24/98	663	RTA00000403F.b.12.1	M00001455D:A06 M00001589C:E06	38985
1419	2/24/98	664	RTA00000404F.a.09.1	M00001389C.E00 M00004039A:C03	9458
1420	2/24/98	665	RTA00000419F.p.20.1	M00004039A:C03	78615
1421	2/24/98	666	RTA00000403F.o.19.1 RTA00000405F.h.07.2	M00001582D:102	4984
1422	2/24/98	667	RTA00000403F.m.07.2 RTA00000408F.m.05.2	M0000157471:G11	23384
1423	2/24/98	668	RTA00000408F.III.05.2 RTA00000410F.b.10.1	M00001530C:B10	74504
1424	2/24/98	669 670	RTA000004101.0.10.11 RTA00000131A.i.6.1	M00001450A:B08	0
1425	2/24/98 2/24/98	670 671	RTA00000131A.i.o.1 RTA00000413F.h.12.1	M00004107A:A12	66929
1426	2/24/98	672	RTA000004151.ii.12.1 RTA00000406F.k.14.1	M00003907C:C02	38651
1427	2/24/98	673	RTA00000406F.d.09.1	M00003875B:F12	38591
1428	2/24/98	674	RTA00000411F.f.17.1	M00003814B:F12	65661
1429	2/24/98	675	RTA00000411F.k.10.1	M00003850D:H11	64506
1430	2/24/98	676	RTA00000411F.g.21.1	M00003823D:G05	64500
1431 1432	2/24/98	677	RTA000004111.g.21.1	M00001457A:C05	82266
1432	1/28/98	412	RTA0000011974.ii.2 iii RTA00000195AF.c.24.1	M00003860D:H07	0
1433	2/24/98	678	RTA00000195AF.c.24.1	M00003860D:H07	0
ננדו	414110	0,0	162		

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln	RTA00000408F.m.22.2	M00001539A:C12	72949
1434	2/24/98	679	RTA00000408F.m.22.2 RTA00000345F.e.11.1	M00001339A:C12 M00001391C:C04	4392
1435	2/24/98	680		M00001391C:C04 M00001464A:D03	34278
1436	2/24/98	681	RTA00000120A.c.24.1	M00001404A:D03	78147
1437	2/24/98	682	RTA00000410F.i.17.1	M00001041B:B01 M00001540D:E02	24723
1438	2/24/98	683	RTA00000403F.j.21.1	M00001340D:E02 M00001426D:D12	6662
1439	2/24/98	684	RTA00000339F.k.20.1	M00001420D.D12 M00001582A:A03	79780
1440	2/24/98	685	RTA00000129A.a.13.2	M00001382A.A03 M00001601A:E12	82067
1441	2/24/98	686	RTA00000129A.k.21.1	M00001601A:E12	9026
1442	2/24/98	687	RTA00000350R.g.10.1		66030
1443	2/24/98	688	RTA00000413F.d.23.1	M00004090B:H06 M00004035A:G10	1937
1444	2/24/98	689	RTA00000419F.p.03.1		0
1445	2/24/98	690	RTA00000341F.b.05.1	M00003793D:A11	8802
1446	2/24/98	691	RTA00000354R.n.08.1	M00003835A:A09	76445
1447	2/24/98	692	RTA00000411F.d.10.1	M00001681D:C12	39281
1448	2/24/98	693	RTA00000404F.b.19.1	M00001592B:A04	
1449	2/24/98	694	RTA00000418F.c.07.1	M00001529D:C05	73245
1450	2/24/98	695	RTA00000418F.j.15.1	M00001632C:H07	74855
1451	2/24/98	696	RTA00000404F.p.12.2	M00001653B:C06	0
1452	2/24/98	697	RTA00000412F.d.14.1	M00003905D:C08	76757
1453	2/24/98	698	RTA00000413F.b.16.1	M00004078A:E05	65126
1454	2/24/98	699	RTA00000340F.I.05.1	M00001644B:D06	38935
1455	2/24/98	700	RTA00000350R.m.14.1	M00001644C:B07	39171
1456	2/24/98	701	RTA00000418F.l.11.1	M00001641C:H07	77158
1457	2/24/98	702	RTA00000130A.d.5.1	M00001605A:H03	82051
1458	2/24/98	703	RTA00000339F.n.05.1	M00001449D:B01	39648
1459	2/24/98	704	RTA00000355R.a.12.1	M00004159C:F09	36756
1460	2/24/98	705	RTA00000407F.a.23.1	M00004081C:A10	23489
1461	2/24/98	706	RTA00000403F.a.09.1	M00001448B:H05	77820
1462	2/24/98	707	RTA00000403F.h.11.1	M00001483B:D04	39219
1463	2/24/98	708	RTA00000406F.j.13.1	M00003905D:B08	38688
1464	2/24/98	709	RTA00000352R.p.09.1	M00004228C:H03	16915
1465	2/24/98	710	RTA00000413F.g.24.1	M00004104D:A04	65481
1466	2/24/98	711	RTA00000404F.1.03.2	M00001636B:G11	40272
1467	2/24/98	712	RTA00000407F.b.18.1	M00004102C:D09	37569
1468	2/24/98	713	RTA00000414F.b.10.1	M00005212D:D09	0
1469	2/24/98	714	RTA00000420F.a.08.1	M00004073A:D10	19473
1470	2/24/98	715	RTA00000418F.b.01.1	M00001475C:G11	76040
1471	2/24/98	716	RTA00000420F.1.03.2	M00005217D:F12	0
1472	2/24/98	717	RTA00000404F.i.22.1	M00001625C:G05	39082
1473	2/24/98	718	RTA00000124A.k.23.1	M00001538A:D03	81350
1474	2/24/98	719	RTA00000404F.e.11.1	M00001608C:E11	38991
1475	2/24/98	720	RTA00000129A.d.2.4	M00001587A:G06	80119
1476	2/24/98	721	RTA00000422F.k.14.1	M00001649D:A08	0
1477	2/24/98	722	RTA00000411F.1.22.1	M00003858B:G05	64439
1478	2/24/98	723	RTA00000419F.o.15.1	M00003989C:D03	32487
1479	2/24/98	724	RTA00000119A.m.17.1	M00001461A:F05	79536
1480	2/24/98	725	RTA00000410F.b.07.1	M00001633C:A05	78916
1481	2/24/98	726	RTA00000420F.b.19.1	M00004088D:A11	36873
1482	2/24/98	727	RTA00000414F.d.02.1	M00005229B:H06	0
1483	2/24/98	728	RTA00000411F.b.21.1	M00001677B:A02	10051

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln	DT 4 00000 402F 20.1	1400001 <i>57(</i> 1.F11	707
1484	2/24/98	729	RTA00000403F.m.20.1	M00001576A:F11	707 16915
1485	2/24/98	730	RTA00000356R.c.16.1	M00004294C:C08	0
1486	2/24/98	731	RTA00000119A.d.17.1	M00001453A:B01	-
1487	2/24/98	732	RTA00000412F.h.11.1	M00003974B:B11	63175
1488	2/24/98	733	RTA00000405F.d.18.1	M00001662C:B02	10494
1489	2/24/98	734	RTA00000414F.e.09.1	M00005236A:G10	0
1490	2/24/98	735	RTA00000420F.a.11.1	M00004073C:D04	66460
1491	2/24/98	736	RTA00000120A.c.7.1	M00001462A:D03	80985
1492	2/24/98	737	RTA00000404F.e.15.1	M00001609B:C09	39101
1493	2/24/98	738	RTA00000422F.n.20.1	M00001669B:B12	38676
1494	2/24/98	739	RTA00000423F.h.20.1	M00003914A:G06	38639
1495	2/24/98	740	RTA00000399F.l.19.1	M00001590D:G07	40145
1496	2/24/98	741	RTA00000414F.b.12.1	M00005212D:H01	0
1497	2/24/98	742	RTA00000410F.b.18.1	M00001633C:H11	76701
1498	2/24/98	743	RTA00000345F.i.08.1	M00001449D:G10	0
1499	2/24/98	744	RTA00000423F.g.15.1	M00003905A:F09	35173
1500	2/24/98	745	RTA00000413F.b.04.1	M00004076D:H07	66427
1501	2/24/98	746	RTA00000345F.e.02.1	M00001395A:E03	0
1502	2/24/98	747	RTA00000413F.n.24.1	M00004960C:E10	0
1503	2/24/98	748	RTA00000346F.f.11.1	M00003793C:D09	38528
1504	2/24/98	749	RTA00000351R.i.13.1	M00003858D:F12	0
1505	2/24/98	750	RTA00000403F.c.05.1	M00001456C:C11	74935
1506	2/24/98	751	RTA00000422F.i.02.1	M00001456C:B12	76436
1507	2/24/98	752	RTA00000410F.a.08.1	M00001632A:B10	73324
1508	2/24/98	753	RTA00000345F.o.13.1	M00001546B:F12	11500
1509	2/24/98	754	RTA00000419F.e.02.1	M00003830C:A03	65010
1510	2/24/98	755	RTA00000423F.d.17.1	M00001663A:C11	20630
1511	2/24/98	756	RTA00000403F.g.13.1	M00001481B:D09	38718
1512	2/24/98	757	RTA00000423F.h.13.1	M00003871A:B09	14398
1513	2/24/98	758	RTA00000407F.a.01.1	M00004039A:H11	12501
1514	2/24/98	759	RTA00000399F.o.06.1	M00001595D:G03	13574
1515	2/24/98	760	RTA00000423F.d.04.1	M00001694A:B12	11307
1516	2/24/98	761	RTA00000411F.f.14.1	M00003814B:C12	62984
1517	2/24/98	762	RTA00000411F.c.04.1	M00001677B:E06	76858
1518	2/24/98	763	RTA00000135A.m.18.1	M00001545A:C03	19255
1519	2/24/98	764	RTA00000413F.c.17.1	M00004085B:B05	36831
1520	2/24/98	765	RTA00000137A.j.15.4	M00001559A:C08	4213
1521	2/24/98	766	RTA00000404F.j.01.1	M00001625D:G10	26859
1522	2/24/98	767	RTA00000138A.p.10.1	M00001644A:H01	81625
1523	2/24/98	768	RTA00000121A.k.5.1	M00001507A:E04	17530
1524	2/24/98	769	RTA00000340F.i.10.1	M00001618A:F10	38561
1525	2/24/98	770	RTA00000421F.f.05.1	M00001477B:E02	5266
1526	2/24/98	771	RTA00000423F.h.07.1	M00003911B:F08	37933
1527	2/24/98	772	RTA00000413F.e.04.1	M00004090C:C07	64176
1528	2/24/98	773	RTA00000406F.h.03.1	M00003901B:A09	38585
1529	2/24/98	774	RTA00000403F.e.24.1	M00001476B:D10	16432
1530	2/24/98	775	RTA00000405F.c.22.1	M00001660C:B06	39053
1531	2/24/98	776	RTA00000403F.i.11.1	M00001485D:E05	23535
1532	2/24/98	777	RTA00000419F.g.02.1	M00003842A:A03	62839
1533	2/24/98	778	RTA00000347F.e.05.1	M00001578D:C04	39814

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
140.	Priority	Priority			•••
	Appln	Appin			
1534	2/24/98	779	RTA00000408F.1.16.1	M00001530A:F12	73468
1535	2/24/98	780	RTA00000405F.I.11.1	M00001693D:E08	2055
1536	2/24/98	781	RTA00000423F.f.09.1	M00003808C:A05	64823
1537	2/24/98	782	RTA00000419F.k.03.1	M00003871C:B05	40822
1538	2/24/98	783	RTA00000406F.b.02.1	M00003867B:G08	38744
1539	2/24/98	784	RTA00000418F.o.14.1	M00001661B:B05	33524
1540	2/24/98	785	RTA00000404F.I.03.1	M00001636B:G11	40272
1541	2/24/98	786	RTA00000404F.b.09.1	M00001591D:C07	39166
1542	2/24/98	787	RTA00000345F.i.24.1	M00001449C:C05	0
1543	2/24/98	788	RTA00000419F.i.04.1	M00003860B:F11	65791
1544	2/24/98	789	RTA00000423F.b.13.1	M00001676C:E07	20619
1545	2/24/98	790	RTA00000345F.n.08.1	M00001517A:B11	0
1546	2/24/98	791	RTA00000399F.n.15.1	M00001594D:C03	3213
1547	2/24/98	792	RTA00000406F.k.11.1	M00003907B:D05	38715
1548	2/24/98	793	RTA00000414F.e.21.1	M00005257C:G01	0
1549	2/24/98	794	RTA00000406F.c.06.1	M00003870C:A01	37924
1550	2/24/98	795	RTA00000418F.n.07.1	M00001658B:A07	76316
1551	2/24/98	796	RTA00000419F.n.15.1	M00003977D:D04	63484
1552	2/24/98	797	RTA00000408F.n.06.2	M00001539A:H12	76642
1553	2/24/98	798	RTA00000420F.c.04.1	M00004089A:B08	65007
1554	2/24/98	799	RTA00000411F.j.15.1	M00003843A:E04	66871
1555	2/24/98	800	RTA00000403F.m.12.1	M00001575D:A02	16933
1556	2/24/98	801	RTA00000128A.m.23.1	M00001561A:D01	81441
1557	2/24/98	802	RTA00000406F.g.03.1	M00003880B:D11	38690
1558	2/24/98	803	RTA00000405F.h.05.2	M00001674A:G07	75706
1559	2/24/98	804	RTA00000129A.n.24.1	M00001604A:C07	81409
1560	2/24/98	805	RTA00000406F.j.08.1	M00003905B:C06	6688
1561	2/24/98	806	RTA00000345F.f.08.1	M00001413B:H09	0
1562	2/24/98	807	RTA00000418F.n.11.1	M00001658D:G12	78977
1563	2/24/98	808	RTA00000418F.p.08.1	M00001669D:D06	73983
1564	2/24/98	809	RTA00000420F.i.23.1	M00005134A:D11	0
1565	2/24/98	810	RTA00000120A.h.9.1	M00001465A:B12	80736
1566	2/24/98	811	RTA00000413F.a.12.1	M00004072D:F09	63403
1567	2/24/98	812	RTA00000412F.o.05.1	M00004034A:A01	63575
1568	2/24/98	813	RTA00000346F.o.06.1	M00004136D:B02	4937
1569	2/24/98	814	RTA00000408F.1.24.1	M00001530B:G09	34263
1570	2/24/98	815	RTA00000403F.a.17.1	M00001448D:E12	13686
1571	2/24/98	816	RTA00000354R.n.04.1	M00003808C:B05	22049
1572	2/24/98	817	RTA00000420F.1.08.2	M00005228C:C05	0
1573	2/24/98	818	RTA00000406F.h.05.1	M00003901B:C03	38542
1574	2/24/98	819	RTA00000410F.b.24.1	M00001633D:D09	75104
1575	2/24/98	820	RTA00000423F.d.11.1	M00001678C:C06	38950
1576	2/24/98	821	RTA00000420F.h.16.1	M00004927A:E06	0
1577	2/24/98	822	RTA00000419F.o.21.1	M00004031A:E01	10336
1578	2/24/98	823	RTA00000119A.k.1.1	M00001460A:H11	81282
1579	2/24/98	824	RTA00000420F.f.07.1	M00004119A:C09	66312
1580	2/24/98	825	RTA00000404F.k.22.2	M00001635D:C12	39084
1581	2/24/98	826	RTA00000422F.e.07.1	M00001579C:G05	38964
1582	2/24/98	827	RTA00000410F.f.12.1	M00001637C:E03	73883
1583	2/24/98	828	RTA00000419F.n.05.1	M00003976C:D06	63713

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln		1400000000 TA D10	72106
1584	2/24/98	829	RTA00000411F.m.11.1	M00003867A:D12	73196
1585	2/24/98	830	RTA00000347F.b.08.1	M00001541B:E05	17591
1586	2/24/98	831	RTA00000420F.d.21.1	M00004107B:B04	65313
1587	2/24/98	832	RTA00000403F.o.10.2	M00001579C:G05	38964 0
1588	2/24/98	833	RTA00000420F.j.20.1	M00005140D:C06	0
1589	2/24/98	834	RTA00000407F.b.11.1	M00004090C:C10	65600
1590	2/24/98	835	RTA00000413F.c.10.1	M00004083B:C01 M00001676D:B02	72893
1591	2/24/98	836	RTA00000411F.b.17.1	M00001876D:B02	0
1592	2/24/98	837	RTA00000420F.h.01.1 RTA00000408F.k.19.1	M00004897C:D00 M00001487C:G03	77593
1593	2/24/98	838 839	RTA00000408F.K.19.1 RTA00000414F.b.01.1	M00001487C:G03	0
1594	2/24/98 2/24/98	839 840	RTA00000414F.b.01.1 RTA00000420F.b.20.1	M00003212B:A02 M00004088D:B05	0
1595	2/24/98	840 841	RTA000004201.b.20.1	M00001457A:G12	82593
1596	2/24/98	842	RTA00000119A.i.s.1	M00001497A:G12 M00003982A:B06	1552
1597	2/24/98	843	RTA000004011.ii.23.1 RTA00000418F.g.03.1	M0000370271:B00	78737
1598	2/24/98	844	RTA00000411F.a.09.1	M00001575C:E00	78629
1599 1600	2/24/98	845	RTA000004111.a.09.1	M00001073C:101	1890
1600	2/24/98	846	RTA00000348K.b.04.1	M00001342B:E01	73183
1602	2/24/98	847	RTA000004131.1.11.1	M00003606C:C07	25073
1602	2/24/98	848	RTA00000404F.n.18.2	M000015712:103	37169
1603	2/24/98	849	RTA00000122A.n.16.1	M00001517A:G08	80553
1605	2/24/98	850	RTA00000420F.c.07.1	M00004089A:E02	65555
1606	2/24/98	85 1	RTA00000423F.d.07.1	M00001678B:B12	0
1607	2/24/98	852	RTA00000414F.f.03.1	M00005257D:G07	0
1608	2/24/98	853	RTA00000408F.j.13.2	M00001485B:D10	42275
1609	2/24/98	854	RTA00000345F.a.07.1	M00001338C:E10	0
1610	2/24/98	855	RTA00000423F.a.01.1	M00001659C:F10	39103
1611	2/24/98	856	RTA00000408F.d.02.1	M00001458D:A01	79169
1612	2/24/98	857	RTA00000404F.e.09.1	M00001608B:A09	39121
1613	2/24/98	858	RTA00000341F.e.20.1	M00003891D:B10	67422
1614	2/24/98	859	RTA00000419F.m.22.1	M00003914A:G09	75600
1615	2/24/98	860	RTA00000419F.m.23.1	M00003958B:E11	64263
1616	2/24/98	861	RTA00000419F.b.06.1	M00001694B:B08	76728
1617	2/24/98	862	RTA00000414F.c.07.1	M00005216A:H01	0
1618	2/24/98	863	RTA00000406F.p.08.1	M00004032C:B02	37573
1619	2/24/98	864	RTA00000129A.n.17.1	M00001604A:A09	79811
1620	2/24/98	865	RTA00000414F.c.03.1	M00005216A:D09	0
1621	2/24/98	866	RTA00000407F.b.08.1	M00004088D:B03	37513
1622	2/24/98	867	RTA00000339F.1.21.1	M00001455D:D11	9781
1623	2/24/98	868	RTA00000406F.i.08.1	M00003903C:E12	37946
1624	2/24/98	869	RTA00000403F.h.07.1	M00001482D:H11	26856
1625	2/24/98	870	RTA00000418F.n.24.1	M00001659D:C09	73153
1626	2/24/98	871	RTA00000403F.f.23.1	M00001479C:E01	39223 74394
1627	2/24/98	872	RTA00000409F.1.20.1	M00001615B:G01 M00001641C:F01	74394
1628	2/24/98	873 874	RTA00000418F.I.06.1	M00001641C:F01 M00004300C:H09	7381
1629	2/24/98	874 875	RTA00000346F.o.22.1	M00004300C:H09	79639
1630	2/24/98	875 876	RTA00000129A.k.22.1 RTA00000423F.d.16.1	M00001601A:E02	39173
1631	2/24/98 2/24/98	876 877	RTA00000423F.d.16.1 RTA00000418F.m.22.1	M00001678D:C11	74567
1632	2/24/98	877 878	RTA00000413F.c.12.1	M00001034D.E12	65334
1633	2124190	0/0	K1A000004151.C.12.1	CODICCOOPERATOR	00007

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
1634	2/24/98	879	RTA00000409F.b.19.1	M00001584D:H02	14479
1635	2/24/98	880	RTA00000418F.g.20.1	M00001585B:C03	74626
1636	2/24/98	881	RTA00000413F.d.15.1	M00004088C:E04	64943
1637	2/24/98	882	RTA00000355R.c.03.1	M00004244C:G07	3986
1638	2/24/98	883	RTA00000406F.c.09.1	M00003870C:E10	5671
1639	2/24/98	884	RTA00000412F.c.10.1	M00003903C:C04	76372
1640	2/24/98	885	RTA00000122A.j.17.1	M00001516A:D02	62736
1641	2/24/98	886	RTA00000420F.m.15.1	M00005235B:F10	0
1642	2/24/98	887	RTA00000339F.p.06.1	M00001484A:A10	4880
1643	2/24/98	888	RTA00000339R.c.04.1	M00001362D:H01	1805
1644	2/24/98	889	RTA00000346F.b.16.1	M00001615C:G05	16485
1645	2/24/98	890	RTA00000418F.j.19.1	M00001634D:D02	78399
1646	2/24/98	89 1	RTA00000137A.p.12.1	M00001587A:B01	80614
1647	2/24/98	892	RTA00000339F.m.17.1	M00001453B:H12	20854
1648	2/24/98	893	RTA00000418F.p.10.1	M00001669D:F05	75323
1649	2/24/98	894	RTA00000408F.k.12.1	M00001486B:D07	77246
1650	2/24/98	895	RTA00000137A.j.11.4	M00001559A:A11	79752
1651	2/24/98	896	RTA00000423F.1.20.1	M00004105C:E09	12580
1652	2/24/98	897	RTA00000419F.n.24.1	M00003980A:F04	65995
1653	2/24/98	898	RTA00000418F.l.03.1	M00001641C:C06	79058
1654	2/24/98	899	RTA00000406F.h.10.1	M00003901C:F09	22732
1655	2/24/98	900	RTA00000419F.m.13.1	M00003908A:F12	79052
1656	2/24/98	901	RTA00000418F.j.14.1	M00001632C:B10	32623
1657	2/24/98	902	RTA00000403F.a.10.1	M00001448C:E11	73952
1658	2/24/98	903	RTA00000420F.a.21.1	M00004078B:C11	66241
1659	2/24/98	904	RTA00000127A.e.6.1	M00001553A:E07	5885
1660	2/24/98	905	RTA00000405F.g.21.2	M00001673B:F07	38966
1661	2/24/98	906	RTA00000405F.g.21.1	M00001673B:F07	38966
1662	2/24/98	907	RTA00000419F.m.06.1	M00003906C:D06	75749
1663	2/24/98	908	RTA00000423F.g.03.1	M00003905C:G11	38007
1664	2/24/98	909	RTA00000420F.i.04.1	M00004959D:H12	0
1665	2/24/98	910	RTA00000418F.f.03.1	M00001577B:F10	78911
1666	2/24/98	911	RTA00000406F.p.13.1	M00004034C:G02	8584
1667	2/24/98	912	RTA00000404F.g.13.1	M00001614C:E06	9436
1668	2/24/98	913	RTA00000120A.c.20.1	M00001464A:B07	43235
1669	2/24/98	914	RTA00000138A.m.15.1	M00001624A:A03	41603
1670	2/24/98	915	RTA00000408F.f.14.2	M00001476D:F03	73024
1671	2/24/98	916	RTA00000418F.p.20.1	M00001677D:B07	78023
1672	2/24/98	917	RTA00000423F.e.21.1	M00003806B:G05	66961
1673	2/24/98	918	RTA00000419F.j.22.1	M00003871A:A02	73525
1674	2/24/98	919	RTA00000410F.d.18.1	M00001635D:D05	75458
1675	2/24/98	920	RTA00000403F.b.24.1	M00001456B:G01	78838
1676	2/24/98	921	RTA00000422F.j.02.1	M00001594D:B08	10368
1677	2/24/98	922	RTA00000410F.e.09.1	M00001636A:F08	76093
1678	2/24/98	923	RTA00000126A.d.19.1	M00001548A:G01	79474
1679	2/24/98	924	RTA00000354R.m.02.1	M00003890B:C08	12766
1680	2/24/98	925	RTA00000353R.h.10.1	M00001390C:C11	39498
1681	2/24/98	926	RTA00000399F.k.20.1	M00001585C:D10	3003
1682	2/24/98	927	RTA00000411F.d.21.1	M00001692B:E01	74794
1683	2/24/98	928	RTA00000340F.m.04.1	M00001679B:H07	19406

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
NO.	Priority	Priority			
	Appln	Appln			
1684	2/24/98	929	RTA00000411F.n.09.1	M00003875A:A07	78962
1685	2/24/98	930	RTA00000127A.h.22.2	M00001554A:E04	13155
1686	2/24/98	931	RTA00000420F.e.09.1	M00004108D:E07	66325
1687	2/24/98	932	RTA00000405F.p.03.1	M00003844A:A11	11346
1688	2/24/98	933	RTA00000419F.a.18.1	M00001680A:B02	78484
1689	2/24/98	934	RTA00000414F.e.01.1	M00005233D:H07	0
1690	2/24/98	935	RTA00000420F.i.07.1	M00004960A:B08	0
1691	2/24/98	936	RTA00000121A.n.23.1	M00001511A:G01	26981
1692	2/24/98	937	RTA00000121A.n.15.1	M00001511A:G08	40849
1693	2/24/98	938	RTA00000403F.i.23.1	M00001487B:E10	11364
1694	2/24/98	939	RTA00000405F.a.03.1	M00001654C:E04	39065
1695	2/24/98	940	RTA00000414F.f.17.1	M00005260A:F04	0
1696	2/24/98	941	RTA00000419F.p.08.1	M00004036D:B04	65560
1697	2/24/98	942	RTA00000126A.n.6.2	M00001551A:D04	79917
1698	2/24/98	943	RTA00000413F.c.03.1	M00004081D:H09	64527
1699	2/24/98	944	RTA00000422F.k.24.1	M00001610C:E06	39118
1700	2/24/98	945	RTA00000412F.c.17.1	M00003905A:A06	75620
1700	2/24/98	946	RTA00000414F.b.07.1	M00005212C:D02	0
1701	2/24/98	947	RTA00000347F.g.08.1	M00004096B:F05	23121
1702	2/24/98	948	RTA00000419F.o.06.1	M00003986C:D09	64643
1703	2/24/98	949	RTA00000340R.j.07.1	M00001654C:D05	38954
1704	2/24/98	950	RTA00000423F.j.02.1	M00003903B:C02	· 38617
1705	2/24/98	951	RTA00000419F.c.04.1	M00003815C:D12	63749
1707	2/24/98	952	RTA00000411F.a.01.1	M00001675B:D02	74524
1707	2/24/98	953	RTA00000406F.f.05.1	M00003878C:F06	22961
1700	2/24/98	954	RTA00000410F.n.05.1	M00001662A:C07	77830
1710	2/24/98	955	RTA00000404F.e.06.1	M00001607D:F06	39315
1711	2/24/98	956	RTA00000423F.I.06.1	M00004062A:H06	38136
1712	2/24/98	957	RTA00000411F.c.03.1	M00001677B:B06	79280
1713	2/24/98	958	RTA00000195AF.c.8.1	M00001678B:H01	0
1713	1/28/98	520	RTA00000195AF.c.8.1	M00001678B:H01	0
1714	2/24/98	959	RTA00000340F.g.20.1	M00001609D:G10	4089
1715	2/24/98	960	RTA00000404F.I.19.2	M00001639B:H01	16196
1716	2/24/98	961	RTA00000420F.n.21.2	M00005259B:D12	0
1717	2/24/98	962	RTA00000404F.p.05.2	M00001652D:E09	1896
1718	2/24/98	963	RTA00000405F.1.07.1	M00001693C:E09	38636
1719	2/24/98	964	RTA00000423F.I.15.1	M00004075B:G09	11219
1720	2/24/98	965	RTA00000411F.n.06.1	M00003871D:E11	73886
1721	2/24/98	966	RTA00000422F.k.15.1	M00001594A:G09	19253
1722	2/24/98	967	RTA00000406F.h.16.1	M00003902B:D06	38618
1723	2/24/98	968	RTA00000419F.f.24.1	M00003841B:E06	18717
1724	2/24/98	969	RTA00000411F.d.18.1	M00001692A:G06	76063
1725	2/24/98	970	RTA00000414F.e.15.1	M00005236B:G03	0
1726	2/24/98	971	RTA00000411F.i.11.1	M00003837C:E05	66849
1727	2/24/98	972	RTA00000408F.d.15.1	M00001459B:C11	78467
1728	2/24/98	973	RTA00000339F.b.22.1	M00001373D:B03	6867
1729	2/24/98	974	RTA00000340F.h.07.1	M00001608D:D11	19254
1730	2/24/98	975	RTA00000411F.n.02.1	M00003870B:F04	78049
1731	2/24/98	976	RTA00000419F.b.17.1	M00003808D:D04	63261
1732	2/24/98	977	RTA00000350R.p.12.1	M00001657C:C07	0

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln	RTA00000130A.e.20.1	M00001606A:H09	79502
1733	2/24/98	978	RTA00000130A.e.20.1 RTA00000345F.b.17.1	M00001000A:1109 M00001362C:H11	945
1734	2/24/98	979		M00001302C:F11	66138
1735	2/24/98	980	RTA00000411F.i.13.1	M00003837C:110	64762
1736	2/24/98	981	RTA00000420F.e.20.1	M00004110B:A07	80915
1737	2/24/98	982	RTA00000126A.p.23.2	M00001332A.F00 M00003809A:H04	0
1738	2/24/98	983	RTA00000423F.f.11.1	M00003889C:H03	37963
1739	2/24/98	984	RTA00000406F.g.08.1	•	74978
1740	2/24/98	985	RTA00000409F.a.08.1	M00001582D:B01	37997
1741	2/24/98	986	RTA00000406F.d.24.1	M00003876B:C05	2368
1742	2/24/98	987	RTA00000422F.b.22.1	M00004117B:A12	2368 15570
1743	2/24/98	988	RTA00000407F.a.22.1	M00004081A:G01	78971
1744	2/24/98	989	RTA00000418F.i.12.1	M00001592A:E02	
1745	2/24/98	990	RTA00000121A.h.19.1	M00001471A:D04	80334
1746	2/24/98	991	RTA00000419F.b.10.1	M00001694C:G04	78566
1747	2/24/98	992	RTA00000406F.m.10.1	M00003914D:B02	38004
1748	2/24/98	993	RTA00000406F.o.05.1	M00003985B:G04	37894
1749	2/24/98	994	RTA00000408F.b.04.2	M00001455A:F04	39933
1750	2/24/98	995	RTA00000411F.k.04.1	M00003850D:A05	65407
1751	2/24/98	996	RTA00000423F.j.03.1	M00003903B:D03	5391
1752	2/24/98	997	RTA00000134A.l.9.1	M00001535A:D10	81814
1753	2/24/98	998	RTA00000341F.g.22.1	M00003914D:D10	0
1754	2/24/98	999	RTA00000418F.k.04.1	M00001637A:A03	75864
1755	2/24/98	1000	RTA00000351R.j.21.1	M00003859D:C05	31604
1756	2/24/98	1001	RTA00000413F.p.07.2	M00005102C:D03	0
1757	2/24/98	1002	RTA00000419F.p.18.1	M00004038D:G06	63002
1758	2/24/98	1003	RTA00000420F.k.08.2	M00005176C:C09	0
1759	2/24/98	1004	RTA00000419F.a.24.1	M00001680B:D02	79290
1760	2/24/98	1005	RTA00000339F.e.17.1	M00001397D:G08	7568
1761	2/24/98	1006	RTA00000129A.e.14.1	M00001587A:F08	80053
1762	2/24/98	1007	RTA00000404F.a.01.1	M00001589B:B08	19251
1763	2/24/98	1008	RTA00000414F.f.07.1	M00005259C:B05	0
1764	2/24/98	1009	RTA00000399F.o.24.1	M00001607D:A11	2272
1765	2/24/98	1010	RTA00000408F.n.16.2	M00001540C:B03	73720
1766	2/24/98	1011	RTA00000400F.c.04.1	M00001618A:F08	6445
1767	2/24/98	1012	RTA00000403F.g.06.1	M00001480C:A05	10505
1768	2/24/98	1013	RTA00000404F.b.18.1	M00001592A:H05	13669
1769	2/24/98	1014	RTA00000412F.I.14.1	M00004029B:F01	62792
1770	2/24/98	1015	RTA00000129A.b.6.2	M00001582A:H01	39111
1771	2/24/98	1016	RTA00000406F.n.12.1	M00003960A:G07	37517
1772	2/24/98	1017	RTA00000418F.e.03.1	M00001573B:G08	73442
1773	2/24/98	1018	RTA00000413F.j.21.1	M00004688A:A02	0
1774	2/24/98	1019	RTA00000403F.g.03.1	M00001479D:G06	23537
1775	2/24/98	1020	RTA00000412F.p.06.1	M00004038B:H10	65485
1776	2/24/98	1021	RTA00000419F.b.21.1	M00003809A:F01	65366
1777	2/24/98	1022	RTA00000401F.j.15.1	M00003901A:C09	3061
1778	2/24/98	1023	RTA00000404F.f.12.1	M00001611B:A05	39209
1779	2/24/98	1024	RTA00000351R.j.16.1	M00003857B:F07	64773
1780	2/24/98	1025	RTA00000118A.j.24.1	M00001450A:B03	18
1781	2/24/98	1026	RTA00000419F.f.18.1	M00003839D:E11	64047
1782	2/24/98	1027	RTA00000423F.i.16.1	M00003907D:A12	38604
			1.00		

SEQ ID	Filing	SEQ ID	Sequence Name	Clone Name	Cluster
NO:	Date of	NO: in			ID
	Priority	Priority			
	Appln	Appln			
1783	2/24/98	1028	RTA00000346F.d.12.1	M00001676B:B09	11777
1784	2/24/98	1029	RTA00000411F.f.04.1	M00003813A:G04	64526
1785	2/24/98	1030	RTA00000125A.c.17.1	M00001542A:E04	80619
1786	2/24/98	1031	RTA00000404F.g.08.1	M00001613D:H10	38980
1787	2/24/98	1032	RTA00000423F.c.13.1	M00001678A:A11	39059
1788	2/24/98	1033	RTA00000414F.e.19.1	M00005257C:E05	0
1789	2/24/98	1034	RTA00000124A.f.16.3	M00001536A:F11	47430
1790	2/24/98	1035	RTA00000404F.k.15.1	M00001634A:B04	18225
1791	2/24/98	1036	RTA00000339F.k.08.1	M00001439B:A10	8133
1792	2/24/98	1037	RTA00000339F.1.12.1	M00001450A:G11	7711
1793	2/24/98	1038	RTA00000406F.b.01.1	M00003867B:G07	39006
1794	2/24/98	1039	RTA00000407F.c.08.1	M00004118D:B05	37549
1795	2/24/98	1040	RTA00000348R.o.12.1	M00001433C:F10	2263
1796	2/24/98	1041	RTA00000403F.b.05.1	M00001455B:E07	74300
1797	2/24/98	1042	RTA00000339F.g.10.1	M00001400C:D02	6327
1798	2/24/98	1043	RTA00000423F.b.17.1	M00001662B:F06	8200
1799	2/24/98	1044	RTA00000419F.n.11.1	M00003977C:B03	66477
1800	2/24/98	1045	RTA00000408F.j.05.2	M00001483C:G06	73878
1801	2/24/98	1046	RTA00000346F.j.06.1	M00003879A:A02	5767
1802	2/24/98	1047	RTA00000419F.c.14.1	M00003819B:G01	65727
1803	2/24/98	1048	RTA00000413F.o.07.2	M00005100A:C01	0
1804	2/24/98	1049	RTA00000405F.f.05.1	M00001669C:D09	14359
1805	2/24/98	1050	RTA00000405F.f.05.2	M00001669C:D09	14359
1806	2/24/98	1051	RTA00000346F.h.24.1	M00003797A:C11	4379
1807	2/24/98	1052	RTA00000420F.b.02.1	M00004081A:A08	64013
1808	2/24/98	1053	RTA00000413F.b.24.1	M00004080A:F01	65117
1809	2/24/98	1054	RTA00000412F.d.08.1	M00003905C:B02	75328
1810	2/24/98	1055	RTA00000346F.a.04.1	M00001607B:C05	5382
1811	2/24/98	1056	RTA00000419F.m.18.1	M00003908C:G09	76014
1812	2/24/98	1057	RTA00000419F.J.24.1	M00003904D:B10	74628
1813	2/24/98	1058	RTA00000408F.c.06.1	M00001456D:E08	78619
1814	2/24/98	1059	RTA00000405F.h.21.2	M00001675C:D12	39072
1815	2/24/98	1060	RTA00000346F.g.02.1	M00003792A:B10	6901
1816	2/24/98	1061	RTA00000405F.g.05.2	M00001671D:E10	38987
1817	2/24/98	1062	RTA00000411F.f.20.1	M00003816C:C01	63501
1818	2/24/98	1063	RTA00000132A.n.7.1	M00001466A:F08	0
1819	2/24/98	1064	RTA00000420F.d.19.1	M00004105C:C08	43146
1820	1/28/98	595	RTA00000195R.a.06.1	M00001394A:E04	35265
1820	2/24/98	1065	RTA00000195R.a.06.1	M00001394A:E04	35265
1821	2/24/98	1066	RTA00000123A.f.2.1	M00001531A:H03	80379
1822	2/24/98	1067	RTA00000411F.j.11.1	M00003841D:F06	66154
1823	2/24/98	1068	RTA00000341F.f.03.1	M00003850A:F06	0
1824	2/24/98	1069	RTA00000346F.k.05.1	M00003904C:A08	0
1825	2/24/98	1070	RTA00000346F.n.22.1	M00004137A:D06	0
1826	2/24/98	1071	RTA00000404F.k.18.2	M00001635A:C06	5475
1827	2/24/98	1072	RTA00000419F.j.03.1	M00003868B:G06	77578
1828	2/24/98	1073	RTA00000418F.a.10.1	M00001475B:C04	15245
1829	2/24/98	1074	RTA00000423F.h.11.1	M00003867C:E11	38977
1830	2/24/98	1075	RTA00000413F.b.17.1	M00004078A:F07	21704
1831	2/24/98	1076	RTA00000423F.k.09.1	M00004035B:H09	26630

Priority	SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
1832 2/24/98 1077		Priority	Priority			
1833 2/24/98 1078	1022	• •	• •	DT 4 00000414F - 11 1	M00005226D. A 12	0
1834 224/98 1079 RTA00000419F.e.10.1 M00003833B:B03 63225 1835 22/4/98 1081 RTA00000435F.d.02.1 M0000145B:D01 39224 1837 22/4/98 1082 RTA00000137A.o.22.1 M00001638D:D01 0 1838 2724/98 1083 RTA00000413F.j.20.1 M00001638A:D01 0 77101 1839 2724/98 1084 RTA00000413F.j.22.1 M00001583A:D01 25076 1840 22/4/98 1085 RTA00000403F.n.22.1 M0000158B:B05 26775 1841 2724/98 1086 RTA00000403F.n.22.2 M00001578B:B05 26775 1841 2724/98 1086 RTA00000403F.n.22.1 M00001578B:B05 26775 1841 2724/98 1087 RTA00000403F.n.22.1 M00001578B:B05 26775 1842 2724/98 1088 RTA00000403F.n.22.1 M00001578B:B05 26775 1844 2724/98 1088 RTA00000339R.b.02.1 M000013906A:H07 17822 1845 2724/98 1099 RTA00000405F.g.22.1 M00001673C:A02 527 1846 2724/98 1099 RTA00000405F.g.22.1 M00001673C:A02 527 1846 2724/98 1091 RTA00000356R.h.05.1 M00001407C:C02 35052 1849 2724/98 1092 RTA00000145F.o.23 M00001673C:A02 527 1849 2724/98 1094 RTA00000356R.h.05.1 M00001407C:C02 35052 1849 2724/98 1095 RTA00000140F.i.15.1 M00001629C:E07 26815 1851 2724/98 1095 RTA00000145F.o.03.1 M00001639C:E07 26815 1851 2724/98 1096 RTA00000419F.o.03.1 M0000169DC:F10 76699 RTA00000409F.i.16.1 M00001590C:F10 76699 RTA00000409F.i.16.1 M00001590C:F10 76699 RTA00000409F.i.16.1 M00001590C:F10 76699 RTA00000409F.i.16.1 M00001590C:F10 76699 RTA00000409F.j.12.1 M00001590C:F10 76699 RTA00000409F.j.12.2 M00001639B:C041 3307 RTA00000409F.j.12.2 M00001639B:C041 3307 RTA00000409F.j.12.2 M00001639B:C04 3307 RTA00000409F.j.12.2 M00001639B:C04 3307 RTA00000409F.j.12.2 M00001639B:C04 3307 RTA00000409F.j.13 M00001639B:C06 RTA00000409F.j.13 M00001639B:C06 RTA00000409F.j.13 M00001639B:C06 RTA00000409F.j.13 M00001639B:C06 RTA00000409F.j.13 M00001639B:F07 12005 RTA00000409F.j.13 M00001639B:F07 12005						
1835 2/24/98 1080						
1836						
1837 2/24/98 1082 RTA00000137A.o.22.1 M00001587A:D01 0 1838 2/24/98 1083 RTA00000418F.j.20.1 M00001634D:D04 77101 1839 2/24/98 1084 RTA00000403F.o.22.2 M000015383:D01 25076 1840 2/24/98 1085 RTA00000403F.o.22.1 M00001578B:B05 26775 1841 2/24/98 1086 RTA00000401F.o.13.1 M00004040C:A01 3220 1843 2/24/98 1088 RTA00000393R.b.02.1 M00001344B:F12 0 1844 2/24/98 1089 RTA00000405F.j.21.1 M000013763C:A02 227 1845 2/24/98 1091 RTA00000356R.h.05.1 M00001673C:A02 257 1847 2/24/98 1091 RTA00000340F.j.15.1 M00001673C:A02 257 1846 2/24/98 1092 RTA0000041F.j.21.1 M00001673C:A02 257 1847 2/24/98 1092 RTA0000041F.j.21.1 M00001673D:F10 2633 1850 2/24/98 1093						
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1842 2/24/98 1087 RTA00000401F.o.13.1 M00004040C:A01 3220 1843 2/24/98 1088 RTA00000339R.b.02.1 M00001344B:F12 0 1844 2/24/98 1099 RTA00000406F.j.2.1.1 M000003906A:H07 17822 1846 2/24/98 1090 RTA00000356R.h05.1 M00001673C:A02 527 1847 2/24/98 1092 RTA00000356R.h05.1 M00001673C:A02 5052 1847 2/24/98 1093 RTA00000340F.i.15.1 M00001629C:E07 26815 1849 2/24/98 1093 RTA00000340F.i.15.1 M00001673D:F10 20633 1850 2/24/98 1095 RTA0000043F.e.12.1 M0000153D:F10 20633 1851 2/24/98 1095 RTA0000041F.a.06.1 M00001589C:A11 23824 1852 2/24/98 1097 RTA0000041F.a.06.1 M00001589C:A11 2385 1855 2/24/98 1098 RTA0000040F.m.16.1 M00001590C:F10 76090 1856 2/24/98 1100						
1843 2/24/98 1088 RTA00000339R.b.02.1 M00001344B:F12 0 1844 2/24/98 1089 RTA00000406F.j.21.1 M00003906A:H07 17822 1845 2/24/98 1090 RTA00000405F.g.22.1 M00001673C:A02 527 1846 2/24/98 1091 RTA00000356R.h.05.1 M00001407C:C02 35052 1847 2/24/98 1093 RTA00000125A.c.2.1 M00001629C:E07 20633 1849 2/24/98 1093 RTA00000345F.h.03.2 M00001673D:F10 20633 1850 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 1851 2/24/98 1095 RTA00000412F.o.03.1 M00001890:F10 765039 1853 2/24/98 1097 RTA00000412F.o.03.1 M0000163D:F10 76090 1854 2/24/98 1098 RTA00000414F.a.10.1 M00001590:F10 76090 1855 2/24/98 1100 RTA00000414F.a.12.1 M00001590:F10 76090 1855 2/24/98 1101 </td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>						
1844 2/24/98 1089 RTA00000406F.j.21.1 M00003906A:H07 17822 1845 2/24/98 1090 RTA00000405F.g.22.1 M00001673C:A02 527 1846 2/24/98 1091 RTA00000356R.h.05.1 M000014707:C02 35052 1847 2/24/98 1092 RTA00000340F.i.15.1 M0000152A::F06 40148 1848 2/24/98 1094 RTA00000345F.c.12.1 M00001673D:F10 26631 1850 2/24/98 1095 RTA00000345F.c.12.1 M00001376:C05 23824 1851 2/24/98 1096 RTA00000412F.o.03.1 M00001580C:A11 2385 1852 2/24/98 1096 RTA00000409F.d.l.6.1 M00001580C:A11 2385 1853 2/24/98 1098 RTA00000409F.d.l.6.1 M00001590C:F10 76090 1855 2/24/98 1099 RTA00000409F.d.l.6.1 M00001590C:F10 76090 1855 2/24/98 1100 RTA00000408F.j.l.7.2 M00001485B:H03 78935 1857 2/24/98 <						
1845 2/24/98 1090 RTA00000405F.g.22.1 M00001673C:A02 527 1846 2/24/98 1091 RTA00000356R.h.05.1 M00004107C:C02 35052 1847 2/24/98 1092 RTA00000125A.c.2.1 M00001542A:F06 40148 1848 2/24/98 1093 RTA00000340F.i.15.1 M00001629C:E07 26815 1849 2/24/98 1094 RTA00000349F.e.l2.1 M00001370E:D0 20633 1850 2/24/98 1095 RTA0000049F.h.03.2 M00001370E:D1 20633 1851 2/24/98 1096 RTA00000421F.a.06.1 M00001589C:A11 2385 1852 2/24/98 1097 RTA00000412F.o.03.1 M00001589C:A11 2385 1853 2/24/98 1099 RTA00000409F.m.16.1 M00001589C:F10 76090 1855 2/24/98 1101 RTA0000049F.d.l6.1 M00001589C:F10 76090 1856 2/24/98 1102 RTA0000049F.j.2.1 M00001549A:H11 40425 1857 2/24/98 1102 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>						
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1847 2/24/98 1092 RTA00000125A.c.2.1 M00001542A:F06 40148 1848 2/24/98 1093 RTA00000340F.i.15.1 M00001673D:F10 20633 1849 2/24/98 1094 RTA00000405F.h.03.2 M00001673D:F10 20633 1850 2/24/98 1095 RTA000004345F.c.12.1 M00001376A:C05 23824 1851 2/24/98 1096 RTA0000042F.a.06.1 M00001589C:A11 2385 1852 2/24/98 1097 RTA0000040F.d.16.1 M00001590C:F10 76090 1853 2/24/98 1098 RTA0000040F.m.16.1 M00001590C:F10 76090 1854 2/24/98 1099 RTA0000040F.m.16.1 M00001590C:F10 76090 1855 2/24/98 1100 RTA0000040F.m.16.1 M00001549A:H11 40425 1857 2/24/98 1101 RTA00000446F.a.16.1 M00001549A:H11 40425 1858 2/24/98 1103 RTA0000046F.a.16.1 M00001547A:F06 0 1859 2/24/98 1104 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>						
1848 2/24/98 1093 RTA00000340F.i.15.1 M00001629C:E07 26815 1849 2/24/98 1094 RTA00000405F.h.03.2 M00001673D:F10 20633 1850 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 1851 2/24/98 1096 RTA00000421F.a.06.1 M00001589C:A11 2385 1852 2/24/98 1097 RTA00000412F.o.03.1 M00004033D:D07 65039 1853 2/24/98 1099 RTA00000409F.d.16.1 M00001590C:F10 76090 1854 2/24/98 1100 RTA00000419F.d.16.1 M00001600B:E04 3307 1855 2/24/98 1101 RTA00000408F.j.17.2 M00001485B:H03 78935 1857 2/24/98 1102 RTA0000046F.a.16.1 M00001590A:H11 40425 1858 2/24/98 1103 RTA0000046F.a.16.1 M0000159A:H11 40425 1858 2/24/98 1103 RTA00000346F.a.16.1 M0000159A:H11 40425 1859 2/24/98 1105 RTA00000346F.a.16.1 M00001547A:F06 0 1860 2/24/98 1105 RTA00000340F.p.18.1 M00003751C:A04 287 1861 2/24/98 1106 RTA00000419F.b.17.1 M00001632B:G10 9579 1864 2/24/98 1107 RTA00000419F.b.17.1 M00001632B:G10 9579 1864 2/24/98 1108 RTA00000346F.a.16.1 M00001652B:G10 9579 1866 2/24/98 1101 RTA00000419F.b.22.1 M0000159A:G3:G10 9579 1866 2/24/98 1110 RTA00000419F.c.16.1 M00001652B:G10 9579 1866 2/24/98 1111 RTA00000419F.c.05.1 M00001631D:G05 39067 1866 2/24/98 1111 RTA00000418F.p.21.1 M00001677D:F03 78068 1869 2/24/98 1114 RTA00000419F.c.05.1 M0000187B:F02 76475 1870 2/24/98 1115 RTA00000419F.d.2.1 M0000167D:F03 78068 1869 2/24/98 1114 RTA00000419F.d.2.1 M00001677D:F03 78068 1871 2/24/98 1115 RTA00000419F.d.2.1 M00001859B:F12 1720						
1849 2/24/98 1094 RTA00000405F.h.03.2 M00001673D:F10 20633 1850 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 1851 2/24/98 1096 RTA00000421F.a.06.1 M00001589C.A11 2385 2/24/98 1098 RTA00000409F.d.16.1 M00001589C:F10 76090 1854 2/24/98 1099 RTA00000409F.d.16.1 M00001590C:F10 76090 1854 2/24/98 1100 RTA00000414F.a.12.1 M00005210A:E06 0 RS55 2/24/98 1101 RTA00000414F.a.12.1 M00005210A:E06 0 RS56 2/24/98 1102 RTA00000408F.j.17.2 M0000148B:H03 78935 1857 2/24/98 1102 RTA00000126A.b.10.1 M0000159A:B07 12082 1859 2/24/98 1103 RTA00000346F.a.16.1 M00001593A:B07 12082 1859 2/24/98 1104 RTA00000126A.b.10.1 M00001547A:F06 0 RTA00000346F.a.16.1 M00001547A:F06 0 RTA00000340F.a.12.2.1 M00001547A:F06 0 RTA00000340F.a.12.2.1 M00001547A:F06 0 RTA00000340F.a.12.1 M00001547A:F06 0 RTA00000340F.a.12.1 M00001547A:F06 0 RTA00000340F.a.12.1 M0000163D:C06 78444 RTA00000346F.a.16.1 M00001653C:H05 77458 RS66 2/24/98 1108 RTA00000418F.a.2.1 M00001653C:H05 38038 RS66 2/24/98 1110 RTA00000418F.a.2.1 M00001653D:G05 39067 RS66 2/24/98 1111 RTA00000418F.a.2.1 M00001653D:C012 38038 RS66 2/24/98 1112 RTA00000418F.a.2.1 M00001677D:F03 78068 RS69 2/24/98 1114 RTA00000418F.a.2.1 M00001677D:F03 78068 RS69 2/24/98 1115 RTA00000419F.a.2.1 M00001677D:F03 78068 RS69 2/24/98 1115 RTA00000410F.o.04.1 M0000167D:F04 79018 RS72 2/24/98 1116 RTA00000410F.o.04.1 M0000167D:F04 79018 RS72 2/24/98 1116 RTA00000353R.l.23.1 M0000148F.C01 6263 RS74 2/24/98 1121 RTA00000404P.f.3.1 M00001457A:B03 81376 2/24/98 1122 RTA00000404P.f.3.1 M000001457A:B03 81						
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1872 2/24/98 1117 RTA00000411F.I.16.1 M00003857C:G01 16122 1873 2/24/98 1118 RTA00000411F.j.03.1 M00003841C:F01 66263 1874 2/24/98 1119 RTA00000126A.k.24.1 M00001550A:F07 39428 1875 2/24/98 1120 RTA00000353R.I.23.1 M00001418B:F07 12531 1876 2/24/98 1121 RTA00000120A.m.10.3 M00001467A:B03 81376 1877 2/24/98 1122 RTA00000419F.f.16.1 M00003839D:E02 64679 1878 2/24/98 1123 RTA00000408F.c.23.1 M00001458C:D10 42261 1879 2/24/98 1124 RTA00000123A.h.22.1 M00001532A:C01 17124 1880 2/24/98 1125 RTA00000118A.n.5.1 M00001451A:C10 0 1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
1873 2/24/98 1118 RTA00000411F.j.03.1 M00003841C:F01 66263 1874 2/24/98 1119 RTA00000126A.k.24.1 M00001550A:F07 39428 1875 2/24/98 1120 RTA00000353R.l.23.1 M00001418B:F07 12531 1876 2/24/98 1121 RTA00000120A.m.10.3 M00001467A:B03 81376 1877 2/24/98 1122 RTA00000419F.f.16.1 M00003839D:E02 64679 1878 2/24/98 1123 RTA00000408F.c.23.1 M00001458C:D10 42261 1879 2/24/98 1124 RTA00000123A.h.22.1 M00001532A:C01 17124 1880 2/24/98 1125 RTA00000118A.n.5.1 M00001451A:C10 0 1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
1874 2/24/98 1119 RTA00000126A.k.24.1 M00001550A:F07 39428 1875 2/24/98 1120 RTA00000353R.l.23.1 M00001418B:F07 12531 1876 2/24/98 1121 RTA00000120A.m.10.3 M00001467A:B03 81376 1877 2/24/98 1122 RTA00000419F.f.16.1 M00003839D:E02 64679 1878 2/24/98 1123 RTA00000408F.c.23.1 M00001458C:D10 42261 1879 2/24/98 1124 RTA00000123A.h.22.1 M00001532A:C01 17124 1880 2/24/98 1125 RTA00000118A.n.5.1 M00001451A:C10 0 1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
1875 2/24/98 1120 RTA00000353R.l.23.1 M00001418B:F07 12531 1876 2/24/98 1121 RTA00000120A.m.10.3 M00001467A:B03 81376 1877 2/24/98 1122 RTA00000419F.f.16.1 M00003839D:E02 64679 1878 2/24/98 1123 RTA00000408F.c.23.1 M00001458C:D10 42261 1879 2/24/98 1124 RTA00000123A.h.22.1 M00001532A:C01 17124 1880 2/24/98 1125 RTA00000118A.n.5.1 M00001451A:C10 0 1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
1876 2/24/98 1121 RTA00000120A.m.10.3 M00001467A:B03 81376 1877 2/24/98 1122 RTA00000419F.f.16.1 M00003839D:E02 64679 1878 2/24/98 1123 RTA00000408F.c.23.1 M00001458C:D10 42261 1879 2/24/98 1124 RTA00000123A.h.22.1 M00001532A:C01 17124 1880 2/24/98 1125 RTA00000118A.n.5.1 M00001451A:C10 0 1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
1877 2/24/98 1122 RTA00000419F.f.16.1 M00003839D:E02 64679 1878 2/24/98 1123 RTA00000408F.c.23.1 M00001458C:D10 42261 1879 2/24/98 1124 RTA00000123A.h.22.1 M00001532A:C01 17124 1880 2/24/98 1125 RTA00000118A.n.5.1 M00001451A:C10 0 1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
1878 2/24/98 1123 RTA00000408F.c.23.1 M00001458C:D10 42261 1879 2/24/98 1124 RTA00000123A.h.22.1 M00001532A:C01 17124 1880 2/24/98 1125 RTA00000118A.n.5.1 M00001451A:C10 0 1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
1879 2/24/98 1124 RTA00000123A.h.22.1 M00001532A:C01 17124 1880 2/24/98 1125 RTA00000118A.n.5.1 M00001451A:C10 0 1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
1880 2/24/98 1125 RTA00000118A.n.5.1 M00001451A:C10 0 1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
1881 2/24/98 1126 RTA00000136A.h.6.1 M00001550A:D09 81620						
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SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			1147
1882	2/24/98	1127	RTA00000401F.g.22.1	M00003871A:G09	1147
1883	2/24/98	1128	RTA00000423F.a.02.3	M00001656B:A08	39210
1884	2/24/98	1129	RTA00000401F.m.07.1	M00003907D:F11	2893
1885	2/24/98	1130	RTA00000354R.p.01.1	M00004104C:H12	0
1886	2/24/98	1131	RTA00000418F.e.20.1	M00001576C:G05	73741
1887	2/24/98	1132	RTA00000119A.c.12.1	M00001453A:D08	4882
1888	2/24/98	1133	RTA00000405F.I.03.1	M00001692D:B01	38580
1889	2/24/98	1134	RTA00000418F.m.02.1	M00001650A:A12	74550
1890	2/24/98	1135	RTA00000346F.o.16.1	M00004358D:C02	176
1891	2/24/98	1136	RTA00000406F.c.05.1	M00003870A:H01	22077
1892	2/24/98	1137	RTA00000345F.d.03.1	M00001376B:A08	19230
1893	2/24/98	1138	RTA00000411F.k.21.1	M00003854B:D04	65349
1894	2/24/98	1139	RTA00000404F.h.20.1	M00001619B:A09	15564
1895	2/24/98	1140	RTA00000339F.c.05.1	M00001365A:H10	3908
1896	2/24/98	1141	RTA00000347F.f.08.1	M00003972D:H02	5948
1897	2/24/98	1142	RTA00000418F.i.06.1	M00001591B:B06	75151
1898	2/24/98	1143	RTA00000423F.a.03.1	M00001656B:D05	26796
1899	2/24/98	1144	RTA00000345F.j.09.1	M00001451B:F01	13
1900	2/24/98	1145	RTA00000423F.k.21.2	M00003984D:B08	37499
1901	2/24/98	1146	RTA00000347F.h.02.1	M00004072D:H12	562
1902	2/24/98	1147	RTA00000404F.c.18.1	M00001594A:C01	38982
1903	2/24/98	1148	RTA00000345F.d.23.1	M00001390D:E03	5862
1904	2/24/98	1149	RTA00000339F.b.02.1	M00001344B:F12	0
1905	2/24/98	1150	RTA00000411F.g.24.1	M00003825B:B11	65233
1906	2/24/98	1151	RTA00000405F.g.18.2	M00001672D:E08	5255
1907	2/24/98	1152	RTA00000405F.m.07.1	M00003809B:B02	37733
1908	2/24/98	1153	RTA00000411F.j.07.1	M00003841C:H11	66963
1909	2/24/98	1154	RTA00000403F.m.09.2	M00001575B:G01	26814
1910	2/24/98	1155	RTA00000353R.h.04.1	M00001375B:C06	17123
1911	2/24/98	1156	RTA00000408F.f.10.2	M00001476D:C05	75309
1912	2/24/98	1157	RTA00000422F.m.18.1	M00001647B:E04	23829
1913	2/24/98	1158	RTA00000405F.o.03.1	M00003829C:H05	37575
1914	2/24/98	1159	RTA00000413F.b.18.1	M00004078C:F04	39873
1915	2/24/98	1160	RTA00000400F.g.02.1	M00001638B:E03	1508
1916	2/24/98	1161	RTA00000346F.m.05.1	M00003983B:C08	5644
1917	2/24/98	1162	RTA00000408F.c.10.1	M00001458A:A11	18247
1918	2/24/98	1163	RTA00000341F.b.14.1	M00003763A:C01	5992
1919	2/24/98	- 1164	RTA00000405F.m.21.1	M00003815C:C06	24218
1920	2/24/98	1165	RTA00000408F.c.08.1	M00001456D:G11	73473
1921	2/24/98	1166	RTA00000347F.h.01.1	M00004040A:G12	12043
1922	2/24/98	1167	RTA00000410F.c.06.1	M00001633D:H06	77784
1923	2/24/98	1168	RTA00000421F.b.06.1	M00001567A:B09	2113
1924	2/24/98	1169	RTA00000405F.b.08.1	M00001656B:E01	39182
1925	2/24/98	1170	RTA00000409F.1.24.1	M00001616C:A02	73174
1926	2/24/98	1171	RTA00000406F.j.06.1	M00003905A:F10	38952
1927	2/24/98	1172	RTA00000423F.h.03.1	M00003875D:D09	37903
1928	2/24/98	1173	RTA00000339R.b.07.1	M00001360A:G10	6826
1929	2/24/98	1174	RTA00000121A.k.22.1	M00001507A:C05	79523
1930	2/24/98	1175	RTA00000414F.b.04.1	M00005212B:E01	0
1931	2/24/98	1176	RTA00000411F.m.06.1	M00003858D:G06	24195
			150		

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
1932	2/24/98	1177	RTA00000126A.b.9.1	M00001547A:F11	81279
1933	2/24/98	1178	RTA00000400F.f.11.1	M00001636A:E07	4088
1934	2/24/98	1179	RTA00000341F.o.12.1	M00004144A:F04	2883
1935	2/24/98	1180	RTA00000404F.1.05.1	M00001636D:F09	38671
1936	2/24/98	1181	RTA00000346F.f.14.1	M00003800B:F03	16998
1937	2/24/98	1182	RTA00000346F.d.21.1	M00001670B:G12	6641
1938	2/24/98	1183	RTA00000346F.j.21.1	M00003879D:A08	3095
1939	2/24/98	1184	RTA00000345F.h.08.1	M00001419D:C10	11393
1940	2/24/98	1185	RTA00000413F.b.20.1	M00004079D:G08	66063
1941	2/24/98	1186	RTA00000419F.p.10.1	M00004036D:B09	41448
1942	2/24/98	1187	RTA00000120A.c.19.1	M00001464A:B03	81016
1943	2/24/98	1188	RTA00000341F.o.18.1	M00004169D:B11	37189
1944	2/24/98	1189	RTA00000339F.o.18.1	M00001469B:B01	6641
1945	2/24/98	1190	RTA00000405F.g.02.2	M00001671B:G05	10567
1946	2/24/98	1191	RTA00000340F.i.05.1	M00001614B:E08	0
1947	2/24/98	1192	RTA00000406F.m.17.1	M00003918A:F09	0
1948	2/24/98	1193	RTA00000411F.k.14.1	M00003851A:C10	63987
1949	2/24/98	1194	RTA00000420F.e.05.1	M00004107D:E12	63908
1950	2/24/98	1195	RTA00000422F.e.23.1	M00001567D:B03	19246
1951	2/24/98	1196	RTA00000413F.l.18.1	M00004895D:G07	0
1952	2/24/98	1197	RTA00000128A.j.10.1	M00001560A:H06	80085
1953	2/24/98	1198	RTA00000412F.f.10.2	M00003959A:A03	65405
1954	2/24/98	1199	RTA00000401F.j.23.1	M00003901C:D03	570
1955	2/24/98	1200	RTA00000422F.k.17.1	M00001652A:A01	38955
1956	2/24/98	1201	RTA00000409F.m.02.1	M00001616C:A11	9157
1957	2/24/98	1202	RTA00000347F.h.10.1	M00004206A:E02	22779
1958	2/24/98	1203	RTA00000413F.e.10.1	M00004092C:B03	31033
1959	2/24/98	1204	RTA00000419F.I.02.1	M00003879A:C01	75736
1960	2/24/98	1205	RTA00000419F.k.05.1	M00003871C:E04	11757
1961	2/24/98	1206	RTA00000418F.b.20.1	M00001484D:G05	73560
1962	2/24/98	1207	RTA00000401F.j.21.1	M00003901B:F10	0
1963	2/24/98	1208	RTA00000347F.e.24.1	M00003823B:F07	8188
1964	2/24/98	1209	RTA00000408F.n.05.2	M00001539A:H02	77883
1965	2/24/98	- 1210	RTA00000419F.o.09.1	M00003987B:F08	66396
1966	2/24/98	1211	RTA00000399F.f.14.1	M00001487D:C11	11483
1967	2/24/98	1212	RTA00000349R.o.03.1	M00001551D:H07	23006
1968	2/24/98	1213	RTA00000135A.a.23.1	M00001537A:H05	27054
1969	2/24/98	1214	RTA00000339F.j.07.1	M00001428D:B10	5673
1970	2/24/98	1215	RTA00000422F.o.08.2	M00001659D:D03	26832
1971	2/24/98	1216	RTA00000404F.e.07.1	M00001608A:D03	9034
1972	2/24/98	1217	RTA00000410F.j.17.1	M00001642D:F02	72912
1973	2/24/98	1218	RTA00000418F.m.18.1	M00001653B:G10	76479
1974	2/24/98	1219	RTA00000347F.e.20.1	M00003771B:E05	39911
1975	2/24/98	1220	RTA00000419F.e.23.1	M00003834B:G04	65772
1976	2/24/98	1221	RTA00000403F.o.17.1	M00001582D:A02	23085
1977	2/24/98	1222	RTA00000423F.e.13.1	M00003848A:C09	10998
1978	2/24/98	1223	RTA00000347F.a.14.1	M00001429D:F11	7421
1979	2/24/98	1224	RTA00000122A.h.24.1	M00001514A:A12	48
1980	2/24/98	1225	RTA00000346F.j.13.1	M00003841C:E04	5337
1981	2/24/98	1226	RTA00000414F.c.12.1	M00005218A:F09	0

ano in	D.11	are in	C N	Ol Name	C14
SEQ ID	Filing	SEQ ID	Sequence Name	Clone Name	Cluster
NO:	Date of	NO: in			ID
	Priority	Priority			
1000	Appln	Appln	DT 4 00000 411 F - 05 1	MAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	64664
1982	2/24/98	1227	RTA00000411F.g.05.1	M00003822D:B10	37148
1983	2/24/98	1228	RTA00000404F.h.10.1	M00001618A:A03	
1984	2/24/98	1229	RTA00000422F.n.14.1	M00001642C:G02	26787
1985	2/24/98	1230	RTA00000399F.j.14.1	M00001578C:F05	16942
1986	2/24/98	1231	RTA00000120A.m.13.3	M00001467A:C10	80608
1987	2/24/98	1232	RTA00000412F.i.03.1	M00003975D:C06	65617
1988	2/24/98	1233	RTA00000418F.l.02.1	M00001641C:C05	39316
1989	2/24/98	1234	RTA00000352R.c.20.1	M00003982A:B12	7339
1990	2/24/98	1235	RTA00000411F.j.04.1	M00003841C:F03	66219
1991	2/24/98	1236	RTA00000414F.b.06.1	M00005212C:C03	0
1992	2/24/98	1237	RTA00000414F.c.24.1	M00005229B:H04	0
1993	2/24/98	1238	RTA00000420F.g.09.1	M00004895B:E12	0
1994	2/24/98	1239	RTA00000340F.o.22.1	M00001673B:B07	7356
1995	2/24/98	1240	RTA00000404F.a.18.1	M00001590B:B02	36267
1996	2/24/98	1241	RTA00000408F.1.14.1	M00001530A:E10	12001
1997	2/24/98	1242	RTA00000405F.d.10.1	M00001661C:F11	39000
1998	2/24/98	1243	RTA00000404F.j.19.1	M00001630D:H10	0
1999	2/24/98	1244	RTA00000418F.h.23.1	M00001591A:B08	75153
2000	2/24/98	1245	RTA00000422F.k.22.1	M00001592C:E05	4098
2001	2/24/98	1246	RTA00000418F.j.11.1	M00001626C:E04	73853
2002	2/24/98	1247	RTA00000408F.o.13.1	M00001572A:B05	74895
2003	2/24/98	1248	RTA00000419F.o.07.1	M00003986C:E09	14059
2004	2/24/98	1249	RTA00000419F.n.17.1	M00003978D:G04	63186
2005	2/24/98	1250	RTA00000403F.f.15.1	M00001477D:F10	22768
2006	2/24/98	1251	RTA00000408F.d.03.1	M00001458D:A02	22768
2007	2/24/98	1252	RTA00000400F.g.08.1	M00001639A:C11	1275
2008	2/24/98	1253	RTA00000346F.f.02.1	M00003772C:B12	62757
2009	2/24/98	1254	RTA00000341F.p.11.1	M00004159C:G12	0
2010	2/24/98	1255	RTA00000413F.i.21.1	M00004118B:B04	64066
2011	2/24/98	1256	RTA00000401F.k.19.1	M00003903D:D10	799
2012	2/24/98	1257	RTA00000419F.h.21.1	M00003856C:B08	64828
2013	2/24/98	1258	RTA00000403F.p.05.2	M00001583D:B08	24528
2014	2/24/98	1259	RTA00000420F.1.19.2	M00005231A:H04	0
2015	2/24/98	1260	RTA00000422F.f.18.1	M00001583D:B08	24528
2016	2/24/98	1261	RTA00000404F.m.17.2	M00001643B:E05	0
2017	2/24/98	1262	RTA00000122A.h.4.1	M00001514A:G03	33576
2018	2/24/98	1263	RTA00000341F.i.22.1	M00003911A:F10	7825
2019	2/24/98	1264	RTA00000345F.e.13.1	M00001392C:D05	4366
2020	2/24/98	1265	RTA00000340F.d.07.1	M00001532D:A06	0
2021	2/24/98	1266	RTA00000121A.a.2.1	M00001468A:H10	81843
2022	3/24/98	1	RTA00000527F.g.13.1	M00003845D:A04	36035
2023	3/24/98	2	RTA00000523F.d.19.1	M00003824A:A06	26489
2024	3/24/98	3	RTA00000528F.b.23.1	M00001479C:F10	1605
2025	3/24/98	4	RTA00000426F.h.11.1	M00003905B:H05	75479
2026	3/24/98	5	RTA00000426F.p.04.1	M00004029B:H08	34149
2027	3/24/98	6	RTA00000523F.1.10.1	M00005134B:E01	0
2028	3/24/98	7	RTA00000523F.o.20.1	M00005177B:H02	0
2029	3/24/98	8	RTA00000428F.b.06.1	M00005228A:A09	0
2030	3/24/98	9	RTA00000522F.b.22.1	M00001573B:H12	75181
2031	3/24/98	10	RTA00000527F.f.12.1	M00003829D:D12	5945

SEQ ID NO:	Filing Date of	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Priority Appln	Appln			
2032	3/24/98	друш 11	RTA00000427F.I.11.1	M00005139A:F01	0
2032	3/24/98	12	RTA00000522F.a.23.1	M00001570C:A05	38613
2034	3/24/98	13	RTA00000528F.m.16.1	M00003845D:C03	4468
2035	3/24/98	14	RTA00000523F.b.02.1	M00003806C:A06	65163
2036	3/24/98	15	RTA00000425F.j.14.1	M00001639D:C12	73397
2037	3/24/98	16	RTA00000426F.m.22.1	M00003983A:G02	30002
2038	3/24/98	17	RTA00000527F.p.06.1	M00004029B:G10	1292
2039	3/24/98	18	RTA00000522F.e.16.1	M00001590A:C08	75283
2040	3/24/98	19	RTA00000527F.j.02.2	M00003856A:B07	4896
2041	3/24/98	20	RTA00000522F.o.06.1	M00001659D:A09	26860
2042	3/24/98	21	RTA00000523F.h.17.1	M00003852A:B03	65586
2043	3/24/98	22	RTA00000527F.k.15.1	M00003982A:G03	22688
2044	3/24/98	23	RTA00000522F.p.07.1	M00001670A:C11	76888
2045	3/24/98	24	RTA00000522F.n.08.1	M00001656A:D10	76343
2046	3/24/98	25	RTA00000425F.c.06.1	M00001585D:D11	78041
2047	3/24/98	26	RTA00000427F.b.23.1	M00003973D:F08	64297
2048	3/24/98	27	RTA00000527F.p.02.1	M00004029B:A01	36844
2049	3/24/98	28	RTA00000427F.d.08.1	M00003980C:E12	63967
2050	3/24/98	29	RTA00000524F.b.03.1	M00005212A:D10	0
2051	3/24/98	30	RTA00000426F.m.07.1	M00004028A:G03	63504
2052	3/24/98	31	RTA00000427F.c.10.1	M00003976B:E06	65478
2053	3/24/98	32	RTA00000424F.n.14.1	M00001584D:C11	73008
2054	3/24/98	33	RTA00000524F.b.21.1	M00005216C:B09	0
2055	3/24/98	34	RTA00000424F.m.15.1	M00001612D:F06	73759
2056	3/24/98	35	RTA00000426F.f.11.1	M00003823C:B01	63102
2057	3/24/98	36	RTA00000428F.a.16.1	M00005212D:F08	0
2058	3/24/98	37	RTA00000426F.f.20.1	M00003854C:F01	65134
2059	3/24/98	38	RTA00000528F.i.22.1	M00001661D:D05	2478
2060	3/24/98	39	RTA00000527F.c.23.1	M00003822C:A07	37742
2061	3/24/98	40	RTA00000426F.h.23.1	M00003911A:D12	75964
2062	3/24/98	41	RTA00000525F.b.17.1	M00004037B:A04	24715
2063	3/24/98	42	RTA00000527F.i.19.2	M00003853C:C06	38089
2064	3/24/98	43	RTA00000527F.p.07.1	M00004029C:B03	23343
2065	3/24/98	44	RTA00000527F.p.17.1	M00004030C:D12	17223
2066	3/24/98	45	RTA00000528F.m.12.1	M00003842D:F08	5768
2067	3/24/98	46	RTA00000523F.c.09.1	M00003813C:D08	47389
2068	3/24/98	47	RTA00000523F.e.18.1	M00003829D:A11	62898
2069	3/24/98	48	RTA00000527F.k.21.1	M00003982B:H10	36051
2070	3/24/98	49	RTA00000527F.n.22.1	M00004027A:A08	24175
2071	3/24/98	50	RTA00000522F.k.15.1	M00001652D:G06	76866
2072	3/24/98	51	RTA00000522F.n.02.1	M00001655D:E08	74959
2073	3/24/98	52	RTA00000523F.1.07.1	M00004927C:H11	0
2074	3/24/98	53	RTA00000525F.c.17.1	M00004040A:C08	38160
2075	3/24/98	54	RTA00000425F.f.19.1	M00001653D:G07	32635
2076	3/24/98	55	RTA00000528F.e.23.1	M00001593B:D10	19242
2077	3/24/98	56	RTA00000522F.n.16.1	M00001657D:A10	26769
2078	3/24/98	57	RTA00000427F.c.20.1	M00003978A:E01	26527
2079	3/24/98	58	RTA00000527F.k.06.1	M00003981B:B12	12469
2080	3/24/98	59	RTA00000427F.n.14.1	M00004960B:D12	0
2081	3/24/98	60	RTA00000523F.i.06.1	M00003855A:A01	66341

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
2002	Appln	Appln	RTA00000427F.f.21.1	M00004118B:C11	36853
2082	3/24/98	61	RTA00000427F.I.21.1 RTA00000427F.j.19.1	M00004178B:C17	41395
2083	3/24/98	62	RTA00000427F.J.19.1	M000046777A:G12	75691
2084	3/24/98	63	RTA00000322F.b.01.1 RTA00000424F.i.24.1	M00001576C:B02	79101
2085	3/24/98	64	RTA00000424F.1.24.1 RTA00000523F.c.01.1	M00001370A:G00 M00003810A:A02	65710
2086	3/24/98	65		M00003971C:F09	66891
2087	3/24/98	66	RTA00000427F.b.15.1	M00003971C.109	25560
2088	3/24/98	67	RTA00000527F.e.03.1	M00005138B:D12	0
2089	3/24/98	68	RTA00000523F.n.04.1	M00003138B.D12	76535
2090	3/24/98	69	RTA00000522F.j.15.2	M00001031C:G12	38147
2091	3/24/98	70	RTA00000525F.e.07.1	M00004113C.G03	37603
2092	3/24/98	71	RTA00000527F.j.20.2	M00003850D.E00	66701
2093	3/24/98	72 72	RTA00000426F.f.19.1	M00005213C:G01	00701
2094	3/24/98	73	RTA00000524F.b.12.1	M00003213C.G01 M00003825B:F10	486
2095	3/24/98	74	RTA00000527F.d.19.1		64688
2096	3/24/98	75	RTA00000523F.i.22.1	M00003857A:E12	04000
2097	3/24/98	76	RTA00000523F.I.18.1	M00005134D:A06	43213
2098	3/24/98	77	RTA00000425F.i.17.1	M00001633A:F11	43213
2099	3/24/98	78	RTA00000427F.o.05.1	M00004958B:D01	
2100	3/24/98	79	RTA00000523F.1.15.1	M00005134C:E11	0
2101	3/24/98	80	RTA00000425F.p.12.1	M00001638C:G01	73219
2102	3/24/98	8 1	RTA00000427F.j.07.1	M00004105A:B10	64819
2103	3/24/98	8 2	RTA00000523F.h.15.1	M00003851C:F09	65137
2104	3/24/98	83	RTA00000527F.i.05.2	M00003851C:B06	37481
2105	3/24/98	84	RTA00000527F.k.18.1	M00003982B:C10	11332
2106	3/24/98	85	RTA00000427F.m.21.1	M00004900C:E11	0
2107	3/24/98	86	RTA00000523F.k.01.1	M00003966C:F03	41437
2108	3/24/98	8 7	RTA00000425F.j.11.1	M00001637C:H12	76667
2109	3/24/98	88	RTA00000424F.b.22.4	M00001530A:F11	72971
2110	3/24/98	89	RTA00000527F.n.02.1	M00003986C:G11	24190
2111	3/24/98	90	RTA00000525F.a.03.1	M00004031D:F05	36786
2112	3/24/98	91	RTA00000527F.i.21.2	M00003855A:F01	37490
2113	3/24/98	92	RTA00000424F.a.24.4	M00001448D:E11	73951
2114	3/24/98	93	RTA00000522F.k.14.1	M00001652D:G02	74280
2115	3/24/98	94	RTA00000522F.n.05.1	M00001655D:H11	73260
2116	3/24/98	95	RTA00000523F.c.18.1	M00003817C:A10	66179
2117	3/24/98	96	RTA00000523F.b.13.1	M00003809B:A03	66330
2118	3/24/98	97	RTA00000522F.j.14.2	M00001651C:D11	73123
2119	3/24/98	98	RTA00000527F.p.16.1	M00004030C:C02	23798
2120	3/24/98	99	RTA00000425F.c.20.1	M00001626D:A02	73581
2121	3/24/98	100	RTA00000424F.i.21.1	M00001596A:E07	73482
2122	3/24/98	101	RTA00000523F.j.19.1	M00003966B:D02	65910
2123	3/24/98	102	RTA00000522F.g.19.1	M00001595C:A01	78119
2124	3/24/98	103	RTA00000424F.b.22.1	M00001530A:F11	72971
2125	3/24/98	104	RTA00000527F.b.18.1	M00003810D:H09	37469
2126	3/24/98	105	RTA00000526F.d.01.1	M00004104B:A02	4468
2127	3/24/98	106	RTA00000424F.j.14.1	M00001592B:B02	74311
2128	3/24/98	107	RTA00000523F.n.20.1	M00005174D:H02	0
2129	3/24/98	108	RTA00000525F.e.16.1	M00004117B:G01	36837
2130	3/24/98	109	RTA00000424F.a.01.4	M00001575A:D05	43214
2131	3/24/98	110	RTA00000522F.d.08.1	M00001578B:A06	74284

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority Appln			
2132	Appln 3/24/98	111	RTA00000425F.d.08.1	M00001631A:F06	74350
2132	3/24/98	112	RTA00000523F.n.12.1	M00005173C:A02	0
2134	3/24/98	113	RTA00000527F.g.07.1	M00003840C:C02	37488
2135	3/24/98	114	RTA00000524F.a.23.1	M00005211C:E09	0
2136	3/24/98	115	RTA00000525F.b.05.1	M00004034C:F05	21116
2137	3/24/98	116	RTA00000425F.n.05.1	M00001647D:G07	73965
2138	3/24/98	117	RTA00000523F.d.18.1	M00003822B:G01	64072
2139	3/24/98	118	RTA00000525F.a.02.1	M00004031C:H10	37454
2140	3/24/98	119	RTA00000523F.p.06.1	M00005177D:F09	0
2141	3/24/98	120	RTA00000426F.h.09.1	M00003905B:G03	78797
2142	3/24/98	121	RTA00000427F.n.02.1	M00004900D:B10	0
2143	3/24/98	122	RTA00000523F.o.12.1	M00005177A:B06	0
2144	3/24/98	123	RTA00000427F.g.05.1	M00004069C:C08	63138
2145	3/24/98	124	RTA00000424F.m.12.1	M00001586C:H07	77675
2146	3/24/98	125	RTA00000424F.a.01.1	M00001575A:D05	43214
2147	3/24/98	126	RTA00000527F.m.05.1	M00003985A:C01	17240
2148	3/24/98	127	RTA00000523F.n.10.1	M00005140D:G09	0
2149	3/24/98	128	RTA00000428F.c.02.1	M00005229D:H07	0
2150	3/24/98	129	RTA00000527F.p.18.1	M00004030D:B06	31635
2151	3/24/98	130	RTA00000427F.h.12.1	M00004092C:D08	36894
2152	3/24/98	131	RTA00000523F.c.15.1	M00003813D:G06	36935
2153	3/24/98	132	RTA00000427F.k.17.1	M00004101A:F07	64965
2154	3/24/98	133	RTA00000425F.f.04.1	M00001607A:B06	24633
2155	3/24/98	134	RTA00000424F.c.14.3	M00001476D:A09	76614
2156	3/24/98	135	RTA00000522F.k.10.2	M00001652D:B09	77619
2157	3/24/98	136	RTA00000424F.m.22.1	M00001614C:E11	72943
2158	3/24/98	137	RTA00000527F.h.17.1	M00003848D:G02	37799
2159	3/24/98	138	RTA00000527F.c.22.1	M00003822B:G12	37496
2160	3/24/98	139	RTA00000425F.k.22.1	M00001633C:E12	78123
2161	3/24/98	140	RTA00000424F.m.14.1	M00001612D:D12	77491
2162	3/24/98	141	RTA00000522F.k.19.1	M00001653A:A05	32625
2163	3/24/98	142	RTA00000523F.i.18.1	M00003856B:C04	64463
2164	3/24/98	143	RTA00000425F.j.22.1	M00001633B:E03	73882
2165	3/24/98	144	RTA00000527F.g.23.1	M00003846C:F08	37538
2166	3/24/98	145	RTA00000426F.m.24.1	M00003981A:A07	63943
2167	3/24/98	146	RTA00000527F.i.17.2	M00003853B:C08	37539
2168	3/24/98	147	RTA00000425F.d.21.1	M00001631B:H04	78920
2169	3/24/98	148	RTA00000427F.n.18.1	M00004891D:C11	0
2170	3/24/98	149	RTA00000424F.d.04.3	M00001478A:F12	76505
2171	3/24/98	150	RTA00000424F.d.04.1	M00001478A:F12	76505
2172	3/24/98	151	RTA00000427F.c.12.1	M00003976B:H07	66995
2173	3/24/98	152	RTA00000425F.d.07.1	M00001631A:F12	43197
2174	3/24/98	153	RTA00000527F.l.13.1	M00003983C:F10 M00001596C:F09	36904 40823
2175	3/24/98	154	RTA00000522F.h.13.1	M00001596C:F09 M00001609C:A12	75454
2176	3/24/98	155	RTA00000424F.l.19.1 RTA00000525F.b.22.1	M00001609C:A12 M00004037C:D07	16679
2177	3/24/98	156	RTA00000523F.g.10.1	M00004037C.D07 M00003848B:E07	40694
2178	3/24/98	157	RTA00000323F.g.10.1 RTA00000427F.a.06.1	M00003848B.E07 M00004036A:A11	66550
2179	3/24/98 3/24/98	158 159	RTA00000427F.a.00.1 RTA00000525F.c.19.1	M00004030A.ATT	38159
2180	3/24/98	160	RTA00000523F.C.19.1 RTA00000523F.f.06.1	M00004040B:107	62871
2181	3124170	100	K 1 /1000003231 .1.00.1	,4100005055D.1100	020/1

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
2102	Appln 3/24/98	Appln 161	RTA00000424F.h.10.1	M00001485C:G06	72925
2182	3/24/98	162	RTA000004247.ii.10.1 RTA00000522F.a.12.1	M000011032:000 M00001567A:H05	33515
2183	3/24/98	163	RTA00000522F.h.01.1	M000015977:H05	75010
2184		163	RTA00000522F.n.01.1	M00001373C:E03	0
2185	3/24/98	165	RTA00000325F.e.21.1	M00003174D:D02	77203
2186	3/24/98	166	RTA00000423F.f.07.1	M00001827D:D10	62799
2187	3/24/98	160	RTA00000323F.i.07.1 RTA00000424F.i.20.1	M00003833D:1110 M00001596A:D01	44010
2188	3/24/98 3/24/98	168	RTA00000424F.j.12.1	M000015967C:E05	73827
2189		169	RTA00000424F.J.12.1 RTA00000425F.f.05.1	M000015742:D05	24090
2190	3/24/98		RTA00000423F.d.12.1	M00001007A:D10	64888
2191	3/24/98	170	RTA00000523F.d.12.1	M00003822B:B00 M00003829A:F03	62878
2192	3/24/98	171	RTA00000325F.E.10.1 RTA00000425F.f.11.1	M00003627A:103	79275
2193	3/24/98	172	RTA00000425F.I.11.1 RTA00000426F.m.18.1	M00001036C:C04 M00003986D:G07	62974
2194	3/24/98	173	RTA00000424F.b.21.4	M00003380D:G07	24686
2195	3/24/98	174	RTA00000424F.b.21.4 RTA00000528F.d.18.1	M00001530A.B02	2684
2196	3/24/98	175		M00001582C:E01 M00001595B:G07	76536
2197	3/24/98	176	RTA00000522F.g.15.1 RTA00000522F.n.12.1	M00001595B:G07	74117
2198	3/24/98	177		M00001030A:1112 M00005179B:H02	0
2199	3/24/98	178	RTA00000428F.a.12.1	M00003179B.H02 M00001530D:A11	73110
2200	3/24/98	179	RTA00000424F.d.10.3	M00001330D.A11 M00004687A:C03	0
2201	3/24/98	180	RTA00000523F.k.02.1	M00004887A.C03 M00003808A:F09	28736
2202	3/24/98	181	RTA00000523F.b.06.1	M00005214B:A06	0
2203	3/24/98	182	RTA00000524F.b.17.1	M00003214B.A00 M00003813C:H08	23090
2204	3/24/98	183	RTA00000527F.c.04.1	M00005214B:D11	0
2205	3/24/98	184	RTA00000524F.b.18.1	M00003214B.D11 M00003850C:G09	37630
2206	3/24/98	185	RTA00000527F.h.21.1	M00003830C:G09 M00001585D:F03	76042
2207	3/24/98	186	RTA00000425F.c.07.1	M00001383D.F03 M00005231D:H10	0
2208	3/24/98	187	RTA00000428F.b.23.1	M00003231D.H10	7692
2209	3/24/98	188	RTA00000525F.c.15.1 RTA00000424F.d.22.3	M00004040A.A07	76189
2210	3/24/98	189	RTA00000424F.d.22.3 RTA00000523F.h.12.1	M00001448B:G07	65745
2211	3/24/98	190	RTA00000523F.II.12.1 RTA00000522F.g.22.1	M00003831C:D07	77504
2212	3/24/98	191	RTA00000523F.m.02.1	M00001333C:B12	0
2213	3/24/98	192	RTA00000323F.III.02.1	M00005134D:1103	0
2214	3/24/98	193	RTA00000428F.b.12.1 RTA00000522F.j.12.2	M00003251C:B07	74341
2215	3/24/98	194	RTA00000522F.j.12.2 RTA00000523F.i.08.1	M00001051C:A04	65099
2216	3/24/98	195	RTA00000523F.f.12.1	M00003835A:C12	63751
2217	3/24/98	196	RTA00000325F.j.20.1	M00003640A:C10	26760
2218	3/24/98	197 198	RTA00000423F.g.20.1	M00007033B:X12	0
2219	3/24/98		RTA000003231.0.03.1	M00003175D:1104 M00004076D:B09	64572
2220	3/24/98	199 200	RTA000004271.1.24.1 RTA00000527F.a.13.1	M00003805D:E06	37740
2221	3/24/98	201	RTA00000327F.n.17.1	M00003803D:200	0
2222	3/24/98 3/24/98	201	RTA00000528F.j.11.1	M00001669B:C12	1070
2223	3/24/98	202	RTA000003201.j.11.17 RTA00000427F.p.10.2	M00005102C:F09	0
2224		203	RTA00000424F.a.09.4	M00003102C:103	77833
2225	3/24/98 3/24/98	204	RTA00000424F.a.09.4 RTA00000426F.h.12.1	M00001375C:C11	78093
2226		203	RTA000004201.fl.12.1 RTA00000525F.f.07.1	M00003303C:112	37500
2227	3/24/98 3/24/98	207	RTA000003231.1.07.1	M0000417571:700	79211
2228 2229	3/24/98	207	RTA00000424F.m.10.1	M00001576B:E11	34251
2229	3/24/98	208	RTA000004247.m.10.1	M000013602:E00	63011
2231	3/24/98	210	RTA000004271.g.16.1	M00001594D:G11	78221
2231	JI 24170	210	1171000000221.6.00.1	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln	DT - 0000040451 02 1	N400001497C-C00	74447
2232	3/24/98	211	RTA00000424F.h.03.1	M00001487C:G09 M00001613A:D02	74737
2233	3/24/98	212	RTA00000424F.n.06.1	M00001613A:D02 M00003978A:E09	63990
2234	3/24/98	213	RTA00000427F.c.22.1	M00003978A.E09	77666
2235	3/24/98	214	RTA00000424F.k.12.1		76982
2236	3/24/98	215	RTA00000425F.f.02.1	M00001607A:A01 M00004092C:B12	76982 26494
2237	3/24/98	216	RTA00000427F.h.11.1		75631
2238	3/24/98	217	RTA00000425F.j.16.1	M00001639D:F02	64206
2239	3/24/98	218	RTA00000427F.i.19.1	M00004102C:D01	63803
2240	3/24/98	219	RTA00000427F.f.17.1	M00004115A:B12	76366
2241	3/24/98	220	RTA00000522F.o.18.1	M00001669B:H06	66367
2242	3/24/98	221	RTA00000427F.j.22.1	M00004097D:B05	65845
2243	3/24/98	222	RTA00000426F.p.10.1	M00004033D:C05	76834
2244	3/24/98	223	RTA00000522F.m.02.1	M00001654C:G07	
2245	3/24/98	224	RTA00000527F.k.09.1	M00003981C:F05	213
2246	3/24/98	225	RTA00000527F.d.09.1	M00003824A:G11	10848
2247	3/24/98	226	RTA00000425F.e.15.1	M00001608D:F11	75921
2248	3/24/98	227	RTA00000427F.i.11.1	M00004097C:H08	26635
2249	3/24/98	228	RTA00000523F.o.14.1	M00005177A:H09	0
2250	3/24/98	229	RTA00000424F.n.13.1	M00001584D:B06	74942
2251	3/24/98	230	RTA00000424F.g.14.1	M00001572A:B06	74879
2252	3/24/98	231	RTA00000426F.e.17.1	M00003810C:B06	64089
2253	3/24/98	232	RTA00000527F.i.13.2	M00003852B:G04	2924
2254	3/24/98	233	RTA00000426F.f.13.1	M00003851A:A06	65384
2255	3/24/98	234	RTA00000524F.c.16.1	M00005218D:G10	0
2256	3/24/98	235	RTA00000427F.g.19.1	M00004087A:B05	64611
2257	3/24/98	236	RTA00000527F.o.01.1	M00004027A:D06	19088
2258	3/24/98	237	RTA00000522F.c.01.1	M00001576A:C11	74938
2259	3/24/98	238	RTA00000522F.g.17.1	M00001595B:G10	76486
2260	3/24/98	239	RTA00000523F.j.17.1	M00003966B:A04	63610
2261	3/24/98	240	RTA00000522F.n.14.1	M00001657C:C11	73410
2262	3/24/98	241	RTA00000527F.o.12.1	M00004028B:G08	688
2263	3/24/98	242	RTA00000523F.e.20.1	M00003829D:F03	65164
2264	3/24/98	243	RTA00000424F.c.15.3	M00001476D:F12	73533
2265	3/24/98	244	RTA00000426F.p.09.1	M00004033D:B07	66665
2266	3/24/98	245	RTA00000522F.p.09.1	M00001670A:F09	75204
2267	3/24/98	246	RTA00000426F.m.21.1	M00003983A:F06	64915
2268	3/24/98	247	RTA00000425F.j.21.1	M00001633B:B11	77373
2269	3/24/98	248	RTA00000527F.l.14.1	M00003983D:A09	14935
2270	3/24/98	249	RTA00000523F.h.21.1	M00003853B:C10	41440
2271	3/24/98	250	RTA00000427F.h.24.1	M00004091B:H09	65193 40841
2272	3/24/98	251	RTA00000425F.f.24.1	M00001656D:C04	
2273	3/24/98	252	RTA00000425F.m.03.1	M00001642D:G08	76045 63781
2274	3/24/98	253	RTA00000426F.m.08.1	M00004030B:A12	64799
2275	3/24/98	254	RTA00000523F.d.24.1	M00003824D:D08	
2276	3/24/98	255	RTA00000523F.c.14.1	M00003813D:C02	66015 66492
2277	3/24/98	256	RTA00000523F.b.20.1	M00003809C:H07	75149
2278	3/24/98	257	RTA00000522F.h.07.1	M00001595D:C11	37820
2279	3/24/98	258	RTA00000527F.g.10.1	M00003845A:E12	10815
2280	3/24/98	259	RTA00000528F.m.04.1	M00003830D:H11	0
2281	3/24/98	260	RTA00000524F.b.02.1	M00005212A:A02	U

No. Date of No. In Priority Appln Appln Appln Appln 261 RTA00000427F.i.22.1 M00004104D:B05 63199 2283 3/24/98 262 RTA00000425F.e.09.1 M00001590D:B04 21289 2284 3/24/98 264 RTA00000425F.e.09.1 M00001698C.604 7559 2286 3/24/98 265 RTA00000425F.e.09.1 M00001698.60-604 7559 2288 3/24/98 266 RTA00000425F.i.21.1 M00001698B-G01 63652 2287 3/24/98 267 RTA00000425F.i.21.1 M0000163B:B02 75305 2288 3/24/98 268 RTA00000425F.i.21.1 M0000163B:B07 62851 2299 3/24/98 269 RTA00000427F.k.19.1 M00001603B:B07 62851 2299 3/24/98 270 RTA00000425F.i.02.1 M00003814B:C01 41446 2292 3/24/98 271 RTA00000426F.m.02.1 M00003814B:C01 41446 2292 3/24/98 272 RTA00000426F.m.02.1 M00003983A:D01 31016 2293 3/24/98 272 RTA00000426F.m.02.1 M00003983A:D01 31016 2294 3/24/98 273 RTA00000426F.m.02.1 M00003983A:D01 31016 2295 3/24/98 273 RTA00000525F.i.05.1 M0000385IB:D03 2874 2296 3/24/98 275 RTA00000522F.i.21 M00001654C:D10 78806 3/24/98 276 RTA00000522F.i.20.1 M00001654C:D10 75801 3/24/98 278 RTA00000522F.e.08.1 M00001659C:A00 77310 3/24/98 278 RTA00000522F.e.08.1 M00003974D:E01 4738 2299 3/24/98 278 RTA00000522F.e.0.1 M00003974D:E01 4738 2303 3/24/98 288 RTA00000522F.e.20.1 M00001659C:A05 7688 3/24/98 288 RTA00000522F.e.20.1 M00001630C:A08 74048 2303 3/24/98 288 RTA00000522F.e.20.1 M00001630C:A08 74048 2305 3/24/98 288 RTA00000522F.e.20.1 M00001630C:A08 74048 2305 3/24/98 288 RTA00000522F.e.20.1 M00001630C:A08 74048 2305 3/24/98 288 RTA00000522F.e.01 M0000159D:C01 0 2310 3/24/98 298 RTA00000522F.e.01 M00001570C:G04 0 2310 3/24/98 298 RTA00000527F.e.05.1 M00001570C:G04 0 2312 3/24/98 299 RTA00000527F.e.05.1 M00001570C:G04 0 2313 3/24/98 299 RTA00000527F.e.05.1 M00001570C:G04 0 2313 3/24/98 299	SEQ ID	Filing	SEQ ID	Sequence Name	Clone Name	Cluster
Applin	NO:	Date of	NO: in			ID
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2299 3/24/98 278 RTA00000522F.g.21.1 M00001595C:A09 77310 2300 3/24/98 279 RTA00000528F.b.03.1 M00001455A:D10 2078 2301 3/24/98 280 RTA00000527F.k.20.1 M00001595C:A05 77688 2302 3/24/98 281 RTA00000527F.k.20.1 M00003982B:H07 17148 2303 3/24/98 283 RTA00000427F.h.22.1 M00004108C:E01 64547 2304 3/24/98 283 RTA00000425F.k.20.1 M00001633C:A08 74048 2305 3/24/98 285 RTA00000522F.b.07.1 M00001570D:E05 78634 2306 3/24/98 285 RTA00000522F.b.07.1 M00003858B:G02 63672 2308 3/24/98 287 RTA00000426F.g.19.1 M00003156D:E05 78634 2309 3/24/98 288 RTA00000427F.l.04.1 M00005136D:B07 0 2311 3/24/98 289 RTA00000427F.l.03.1 M00005136D:B07 0 2312 3/24/98 291	2297	3/24/98		RTA00000427F.h.19.1	M00004092D:B11	63047
2300 3/24/98 279 RTA00000528F.b.03.1 M00001455A:D10 2078 2301 3/24/98 280 RTA00000522F.g.20.1 M00001595C:A05 77688 2302 3/24/98 281 RTA00000527F.k.20.1 M00001985C:A05 77688 2303 3/24/98 282 RTA00000427F.h.22.1 M00004108C:E01 64547 2304 3/24/98 283 RTA00000524F.b.19.1 M00001633C:A08 74048 2305 3/24/98 284 RTA00000524F.b.19.1 M00001570D:E05 78634 2306 3/24/98 285 RTA00000525F.d.19.1 M0000385B:B002 63672 2308 3/24/98 286 RTA00000525F.d.19.1 M00004114B:D09 36860 2309 3/24/98 287 RTA00000525F.d.19.1 M00003136D:C01 0 2311 3/24/98 289 RTA00000427F.l.04.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000525F.c.02.1 M00005136D:B07 0 2312 3/24/98 291 <td< td=""><td>2298</td><td>3/24/98</td><td>277</td><td></td><td>M00003974D:E01</td><td>47387</td></td<>	2298	3/24/98	277		M00003974D:E01	47387
2301 3/24/98 280 RTA00000522F.g.20.1 M00001595C:A05 77688 2302 3/24/98 281 RTA00000527F.k.20.1 M00003982B:H07 17148 2303 3/24/98 282 RTA00000427F.k.20.1 M0000163C:Ca08 74048 2304 3/24/98 283 RTA00000425F.k.20.1 M0000163C:Ca08 74048 2305 3/24/98 284 RTA00000525F.b.19.1 M00001570D:E05 78634 2306 3/24/98 285 RTA00000525F.b.07.1 M00001570D:E05 78634 2307 3/24/98 286 RTA00000426F.g.19.1 M00003858B:G02 63672 2308 3/24/98 287 RTA00000427F.l.04.1 M000005136D:C01 0 2310 3/24/98 288 RTA00000427F.l.03.1 M00005136D:B07 0 2311 3/24/98 290 RTA00000427F.l.03.1 M00005177C:G04 0 2312 3/24/98 291 RTA000000525F.c.02.1 M00001575B:C01 77976 2314 3/24/98 293	2299	3/24/98	278	RTA00000522F.g.21.1	M00001595C:A09	77310
2302 3/24/98 281 RTA00000527F.k.20.1 M00003982B:H07 17148 2303 3/24/98 282 RTA00000427F.h.22.1 M00004108C:E01 64547 2304 3/24/98 284 RTA000000524F.b.19.1 M00001633C:A08 74048 2305 3/24/98 284 RTA00000524F.b.19.1 M00001570D:E05 78634 2307 3/24/98 286 RTA0000042F.g.19.1 M00001570D:E05 78634 2307 3/24/98 286 RTA0000042F.b.19.1 M00003858B:G02 63672 2308 3/24/98 287 RTA00000525F.d.19.1 M00005136D:C01 0 2310 3/24/98 288 RTA00000427F.d.10.1 M00005136D:C01 0 2311 3/24/98 290 RTA00000427F.d.03.1 M00005136D:B07 0 2313 3/24/98 291 RTA00000523F.o.23.1 M00001577C:G04 0 2313 3/24/98 292 RTA00000424F.a.05.4 M00001575B:C01 77976 2314 3/24/98 293 RTA	2300	3/24/98	279	RTA00000528F.b.03.1	M00001455A:D10	2078
2303 3/24/98 282 RTA00000427F.h.22.1 M00004108C:E01 64547 2304 3/24/98 283 RTA00000425F.k.20.1 M00001633C:A08 74048 2305 3/24/98 284 RTA00000524F.b.19.1 M0000516B:D02 0 2306 3/24/98 285 RTA00000522F.b.07.1 M00001570D:E05 78634 2307 3/24/98 286 RTA00000525F.d.19.1 M00001570D:E05 78634 2308 3/24/98 287 RTA00000525F.d.19.1 M00004114B:D09 36860 2309 3/24/98 288 RTA00000427F.l.04.1 M00005136D:C01 0 2310 3/24/98 289 RTA00000427F.l.03.1 M00005136D:B07 0 2311 3/24/98 290 RTA00000427F.d.03.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000525F.c.02.1 M0000157B:C01 77976 2314 3/24/98 293 RTA00000424F.a.05.1 M00001554B:A01 74691 2317 3/24/98 295 RTA0	2301	3/24/98	280	RTA00000522F.g.20.1	M00001595C:A05	77688
2304 3/24/98 283 RTA00000425F.k.20.1 M00001633C:A08 74048 2305 3/24/98 284 RTA00000524F.b.19.1 M00005216B:D02 0 2306 3/24/98 285 RTA00000522F.b.07.1 M00001570D:E05 78634 2307 3/24/98 286 RTA00000426F.g.19.1 M00003858B:G02 63672 2308 3/24/98 287 RTA00000427F.l.04.1 M00001136D:C01 0 2310 3/24/98 288 RTA00000427F.l.04.1 M00005136D:C01 0 2311 3/24/98 289 RTA00000427F.l.03.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000523F.o.23.1 M00005176:G04 0 2313 3/24/98 292 RTA00000525F.c.02.1 M00001575B:C01 77976 2314 3/24/98 293 RTA00000525F.c.02.1 M00001575B:C01 77976 2316 3/24/98 293 RTA00000525F.c.02.1 M00001654B:A01 74691 2317 3/24/98 295 RTA0000	2302	3/24/98	281	RTA00000527F.k.20.1	M00003982B:H07	17148
2305 3/24/98 284 RTA00000524F.b.19.1 M00005216B:D02 0 2306 3/24/98 285 RTA00000522F.b.07.1 M00001570D:E05 78634 2307 3/24/98 286 RTA00000426F.g.19.1 M00003858B:G02 63672 2308 3/24/98 287 RTA00000525F.d.19.1 M00005136D:C01 0 2310 3/24/98 288 RTA00000427F.l.04.1 M00005136D:C01 0 2311 3/24/98 289 RTA00000427F.l.03.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000427F.l.03.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000523F.o.23.1 M00005177C:G04 0 2313 3/24/98 292 RTA00000525F.c.02.1 M00001575B:C01 77976 2314 3/24/98 293 RTA00000525F.c.02.1 M00001575B:C01 77976 2316 3/24/98 295 RTA00000525F.c.02.1 M00001654B:A01 74691 2317 3/24/98 296 RTA0000052	2303	3/24/98	282	RTA00000427F.h.22.1	M00004108C:E01	64547
2306 3/24/98 285 RTA00000522F.b.07.1 M00001570D:E05 78634 2307 3/24/98 286 RTA00000426F.g.19.1 M00003858B:G02 63672 2308 3/24/98 287 RTA00000525F.d.19.1 M00004114B:D09 36860 2309 3/24/98 288 RTA00000427F.l.04.1 M00005136D:C01 0 2310 3/24/98 289 RTA00000427F.l.03.1 M000093978C:A12 40685 2311 3/24/98 290 RTA00000427F.l.03.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000427F.l.03.1 M00005177C:G04 0 2313 3/24/98 291 RTA00000523F.o.23.1 M00001577B:C01 77976 2314 3/24/98 292 RTA00000525F.c.02.1 M00004038A:E05 14618 2315 3/24/98 293 RTA00000525F.c.02.1 M00001654B:A01 74691 2316 3/24/98 295 RTA00000525F.c.11.5 M00001654B:A01 74691 2319 3/24/98 297 <	2304	3/24/98	283	RTA00000425F.k.20.1	M00001633C:A08	74048
2307 3/24/98 286 RTA00000426F.g.19.1 M00003858B:G02 63672 2308 3/24/98 287 RTA00000525F.d.19.1 M00004114B:D09 36860 2309 3/24/98 288 RTA00000427F.l.04.1 M00005136D:C01 0 2310 3/24/98 289 RTA00000427F.l.03.1 M00005136D:B07 0 2311 3/24/98 290 RTA00000523F.o.23.1 M00005177C:G04 0 2312 3/24/98 291 RTA00000523F.o.23.1 M00001575B:C01 77976 2314 3/24/98 292 RTA00000525F.c.02.1 M00004038A:E05 14618 2315 3/24/98 293 RTA00000525F.c.02.1 M00001654B:A01 74691 2316 3/24/98 294 RTA00000522F.l.15.1 M00001654B:A01 74691 2317 3/24/98 296 RTA00000525F.c.02.1 M00001654B:A01 74691 2318 3/24/98 297 RTA00000525F.c.01.1 M00000439C:E02 37895 2319 3/24/98 298 <t< td=""><td>2305</td><td>3/24/98</td><td>284</td><td>RTA00000524F.b.19.1</td><td>M00005216B:D02</td><td>0</td></t<>	2305	3/24/98	284	RTA00000524F.b.19.1	M00005216B:D02	0
2308 3/24/98 287 RTA00000525F.d.19.1 M00004114B:D09 36860 2309 3/24/98 288 RTA00000427F.l.04.1 M00005136D:C01 0 2310 3/24/98 289 RTA00000427F.d.10.1 M00003978C:A12 40685 2311 3/24/98 290 RTA00000427F.l.03.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000525F.c.23.1 M00005177C:G04 0 2313 3/24/98 292 RTA00000525F.c.02.1 M00001575B:C01 77976 2314 3/24/98 293 RTA00000525F.c.02.1 M00001575B:C01 77976 2315 3/24/98 293 RTA00000525F.c.02.1 M00001575B:C01 77976 2316 3/24/98 294 RTA00000525F.c.02.1 M00001654B:A01 74691 2317 3/24/98 295 RTA00000525F.c.02.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.c.11.1 M00004039C:E02 37895 2319 3/24/98 298 <t< td=""><td>2306</td><td>3/24/98</td><td>285</td><td>RTA00000522F.b.07.1</td><td>M00001570D:E05</td><td>78634</td></t<>	2306	3/24/98	285	RTA00000522F.b.07.1	M00001570D:E05	78634
2309 3/24/98 288 RTA00000427F.l.04.1 M00005136D:C01 0 2310 3/24/98 289 RTA00000427F.d.10.1 M00003978C:A12 40685 2311 3/24/98 290 RTA00000427F.l.03.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000523F.o.23.1 M00005177C:G04 0 2313 3/24/98 292 RTA00000424F.a.05.4 M00001575B:C01 77976 2314 3/24/98 293 RTA00000525F.c.02.1 M00004038A:E05 14618 2315 3/24/98 294 RTA00000525F.c.02.1 M00001575B:C01 77976 2316 3/24/98 295 RTA00000522F.l.15.1 M00001654B:A01 74691 2317 3/24/98 296 RTA00000525F.c.01.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.c.11.1 M00003826B:B04 19015 2320 3/24/98 298 RTA00000527F.e.08.1 M0000157A:A03 75449 2321 3/24/98 300 <td< td=""><td>2307</td><td>3/24/98</td><td>286</td><td>RTA00000426F.g.19.1</td><td>M00003858B:G02</td><td>63672</td></td<>	2307	3/24/98	286	RTA00000426F.g.19.1	M00003858B:G02	63672
2310 3/24/98 289 RTA00000427F.d.10.1 M00003978C:A12 40685 2311 3/24/98 290 RTA00000427F.l.03.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000523F.o.23.1 M00005177C:G04 0 2313 3/24/98 292 RTA00000424F.a.05.4 M00001575B:C01 77976 2314 3/24/98 293 RTA00000525F.c.02.1 M00004038A:E05 14618 2315 3/24/98 294 RTA00000424F.a.05.1 M00001575B:C01 77976 2316 3/24/98 295 RTA00000522F.l.15.1 M00001654B:A01 74691 2317 3/24/98 296 RTA00000525F.e.02.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.e.08.1 M00004039C:E02 37895 2319 3/24/98 298 RTA00000527F.e.08.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000525F.e.14.1 M00001584A:A07 19402 2322 3/24/98 301	2308	3/24/98	287	RTA00000525F.d.19.1	M00004114B:D09	36860
2311 3/24/98 290 RTA00000427F.I.03.1 M00005136D:B07 0 2312 3/24/98 291 RTA00000523F.o.23.1 M00005177C:G04 0 2313 3/24/98 292 RTA00000424F.a.05.4 M00001575B:C01 77976 2314 3/24/98 293 RTA00000525F.c.02.1 M00004038A:E05 14618 2315 3/24/98 294 RTA00000424F.a.05.1 M00001575B:C01 77976 2316 3/24/98 295 RTA00000522F.I.15.1 M00001654B:A01 74691 2317 3/24/98 296 RTA00000525F.c.02.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.c.11.1 M00004039C:E02 37895 2319 3/24/98 298 RTA00000527F.e.08.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000527F.f.18.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00001567A:C11 73662 2324 3/24/98 302	2309	3/24/98	288	RTA00000427F.1.04.1	M00005136D:C01	0
2312 3/24/98 291 RTA00000523F.o.23.1 M00005177C:G04 0 2313 3/24/98 292 RTA00000424F.a.05.4 M00001575B:C01 77976 2314 3/24/98 293 RTA00000525F.c.02.1 M00004038A:E05 14618 2315 3/24/98 294 RTA00000424F.a.05.1 M00001575B:C01 77976 2316 3/24/98 295 RTA00000522F.l.15.1 M00001654B:A01 74691 2317 3/24/98 296 RTA00000425F.e.02.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.c.11.1 M00004039C:E02 37895 2319 3/24/98 298 RTA00000527F.e.08.1 M00003826B:B04 19015 2320 3/24/98 299 RTA00000522F.c.14.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000527F.f.18.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00005100B:H07 0 2324 3/24/98 303	2310	3/24/98	289	RTA00000427F.d.10.1	M00003978C:A12	40685
2313 3/24/98 292 RTA00000424F.a.05.4 M00001575B:C01 77976 2314 3/24/98 293 RTA00000525F.c.02.1 M00004038A:E05 14618 2315 3/24/98 294 RTA00000424F.a.05.1 M00001575B:C01 77976 2316 3/24/98 295 RTA00000522F.l.15.1 M00001654B:A01 74691 2317 3/24/98 296 RTA00000425F.e.02.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.c.11.1 M00004039C:E02 37895 2319 3/24/98 298 RTA00000527F.e.08.1 M00003826B:B04 19015 2320 3/24/98 299 RTA00000527F.e.14.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000527F.f.18.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00005100B:H07 0 2324 3/24/98 302 RTA00000527F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304	2311	3/24/98	290	RTA00000427F.l.03.1	M00005136D:B07	0
2314 3/24/98 293 RTA00000525F.c.02.1 M00004038A:E05 14618 2315 3/24/98 294 RTA00000424F.a.05.1 M00001575B:C01 77976 2316 3/24/98 295 RTA00000522F.l.15.1 M00001654B:A01 74691 2317 3/24/98 296 RTA00000425F.e.02.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.c.11.1 M00004039C:E02 37895 2319 3/24/98 298 RTA00000527F.e.08.1 M00003826B:B04 19015 2320 3/24/98 299 RTA00000527F.e.08.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000527F.c.14.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00003830D:B11 37577 2323 3/24/98 302 RTA00000427F.p.04.2 M00005100B:H07 0 2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304	2312	3/24/98	291	RTA00000523F.o.23.1	M00005177C:G04	0
2315 3/24/98 294 RTA00000424F.a.05.1 M00001575B:C01 77976 2316 3/24/98 295 RTA00000522F.l.15.1 M00001654B:A01 74691 2317 3/24/98 296 RTA00000425F.e.02.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.c.11.1 M00004039C:E02 37895 2319 3/24/98 298 RTA00000527F.e.08.1 M00003826B:B04 19015 2320 3/24/98 299 RTA00000522F.c.14.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000527F.f.18.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00005100B:H07 0 2324 3/24/98 302 RTA00000522F.a.06.1 M00005100B:H07 0 2324 3/24/98 304 RTA00000525F.d.13.1 M00005173D:H02 0 2325 3/24/98 305 RTA00000525F.d.23.1 M00005100B:G11 0 2327 3/24/98 306 RT	2313	3/24/98	292	RTA00000424F.a.05.4	M00001575B:C01	77976
2316 3/24/98 295 RTA00000522F.l.15.1 M00001654B:A01 74691 2317 3/24/98 296 RTA00000425F.e.02.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.c.11.1 M00004039C:E02 37895 2319 3/24/98 298 RTA00000527F.e.08.1 M00003826B:B04 19015 2320 3/24/98 299 RTA00000527F.e.08.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000527F.f.18.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00003830D:B11 37577 2323 3/24/98 302 RTA00000527F.p.04.2 M00005100B:H07 0 2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000527F.p.03.2 M00005100B:G11 0 2329 3/24/98 307 <t< td=""><td>2314</td><td>3/24/98</td><td>293</td><td>RTA00000525F.c.02.1</td><td></td><td></td></t<>	2314	3/24/98	293	RTA00000525F.c.02.1		
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2317 3/24/98 296 RTA00000425F.e.02.1 M00001625C:F10 76143 2318 3/24/98 297 RTA00000525F.c.11.1 M00004039C:E02 37895 2319 3/24/98 298 RTA00000527F.e.08.1 M00003826B:B04 19015 2320 3/24/98 299 RTA00000522F.c.14.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000424F.m.08.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00003830D:B11 37577 2323 3/24/98 302 RTA00000427F.p.04.2 M00005100B:H07 0 2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000527F.p.03.2 M00005100B:G11 0 2329 3/24/98 307 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 <t< td=""><td></td><td></td><td></td><td></td><td></td><td></td></t<>						
2318 3/24/98 297 RTA00000525F.c.11.1 M00004039C:E02 37895 2319 3/24/98 298 RTA00000527F.e.08.1 M00003826B:B04 19015 2320 3/24/98 299 RTA00000522F.c.14.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000424F.m.08.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00003830D:B11 37577 2323 3/24/98 302 RTA00000427F.p.04.2 M00005100B:H07 0 2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304 RTA00000522F.d.13.1 M00004110C:E03 349 2326 3/24/98 305 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000424F.k.23.1 M00001614A:B10 31061 2329 3/24/98 309						
2319 3/24/98 298 RTA00000527F.e.08.1 M00003826B:B04 19015 2320 3/24/98 299 RTA00000522F.c.14.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000424F.m.08.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00003830D:B11 37577 2323 3/24/98 302 RTA00000427F.p.04.2 M00005100B:H07 0 2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304 RTA00000522F.d.13.1 M00004110C:E03 349 2326 3/24/98 305 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000427F.p.03.2 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384				RTA00000525F.c.11.1		
2320 3/24/98 299 RTA00000522F.c.14.1 M00001577A:A03 75449 2321 3/24/98 300 RTA00000424F.m.08.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00003830D:B11 37577 2323 3/24/98 302 RTA00000427F.p.04.2 M00005100B:H07 0 2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304 RTA00000522F.d.13.1 M00004110C:E03 349 2326 3/24/98 305 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384				RTA00000527F.e.08.1		
2321 3/24/98 300 RTA00000424F.m.08.1 M00001584A:A07 19402 2322 3/24/98 301 RTA00000527F.f.18.1 M00003830D:B11 37577 2323 3/24/98 302 RTA00000427F.p.04.2 M00005100B:H07 0 2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304 RTA00000525F.d.13.1 M00004110C:E03 349 2326 3/24/98 305 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384						
2322 3/24/98 301 RTA00000527F.f.18.1 M00003830D:B11 37577 2323 3/24/98 302 RTA00000427F.p.04.2 M00005100B:H07 0 2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304 RTA00000525F.d.13.1 M00004110C:E03 349 2326 3/24/98 305 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384						
2323 3/24/98 302 RTA00000427F.p.04.2 M00005100B:H07 0 2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304 RTA00000525F.d.13.1 M00004110C:E03 349 2326 3/24/98 305 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384						
2324 3/24/98 303 RTA00000522F.a.06.1 M00001567A:C11 73662 2325 3/24/98 304 RTA00000525F.d.13.1 M00004110C:E03 349 2326 3/24/98 305 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384						
2325 3/24/98 304 RTA00000525F.d.13.1 M00004110C:E03 349 2326 3/24/98 305 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384				-		
2326 3/24/98 305 RTA00000523F.n.16.1 M00005173D:H02 0 2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384						
2327 3/24/98 306 RTA00000522F.d.23.1 M00001579D:F02 73868 2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384						
2328 3/24/98 307 RTA00000427F.p.03.2 M00005100B:G11 0 2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384						
2329 3/24/98 308 RTA00000424F.k.23.1 M00001614A:B10 31061 2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384						
2330 3/24/98 309 RTA00000523F.j.10.1 M00003860B:G09 63384						

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2332	3/24/98	311	RTA00000428F.b.02.1	M00005214D:D10	0
2333	3/24/98	312	RTA00000426F.f.17.1	M00003811C:C02	66334
2334	3/24/98	313	RTA00000523F.j.21.1	M00003966C:A12	36925
2335	3/24/98	314	RTA00000522F.e.09.1	M00001589D:A01	32599
2336	3/24/98	315	RTA00000427F.n.19.1	M00004891D:E07	0
2337	3/24/98	316	RTA00000523F.h.16.1	M00003851D:H11	66031
2338	3/24/98	317	RTA00000428F.a.01.1	M00004897D:G05	0
2339	3/24/98	318	RTA00000523F.a.01.1	M00001671C:F11	74923
2340	3/24/98	319	RTA00000523F.p.15.1	M00005178B:H01	0
2341	3/24/98	320	RTA00000427F.j.06.1	M00004102D:B05	63676
2342	3/24/98	321	RTA00000424F.m.04.1	M00001609C:G05	79017
2343	3/24/98	322	RTA00000523F.i.17.1	M00003856B:A12	65779
2344	3/24/98	323	RTA00000524F.c.12.1	M00005218B:D09	0
2345	3/24/98	324	RTA00000523F.o.09.1	M00005176A:C12	0
2346	3/24/98	325	RTA00000525F.c.18.1	M00004040B:C05	24208
2347	3/24/98	326	RTA00000527F.e.09.1	M00003826B:E11	37521
2348	3/24/98	327	RTA00000424F.j.08.1	M00001596B:D09	73972
2349	3/24/98	328	RTA00000523F.n.01.1	M00005137A:E01	0
2350	3/24/98	329	RTA00000527F.c.09.1	M00003817C:G06	64859
2351	3/24/98	330	RTA00000523F.d.23.1	M00003824C:A10	63633
2352	3/24/98	331	RTA00000528F.k.10.1	M00001678C:F09	1981
2353	3/24/98	332	RTA00000523F.c.03.1	M00003810B:B11	36913
2354	3/24/98	333	RTA00000427F.k.21.1	M00004090D:F12	62880
2355	3/24/98	334	RTA00000427F.n.11.1	M00004960B:A09	0
2356	3/24/98	335	RTA00000427F.d.09.1	M00003980C:F12	66486
2357	3/24/98	336	RTA00000426F.n.17.1	M00004039D:B10	66572
2358	3/24/98	337	RTA00000525F.e.08.1	M00004115C:H04	24193
2359	3/24/98	338	RTA00000523F.e.15.1	M00003829C:E08	7919
2360	3/24/98	339	RTA00000426F.m.03.1	M00004034C:E08	66480
2361	3/24/98	340	RTA00000424F.h.06.1	M00001485C:D07	77552
2362	3/24/98	341	RTA00000425F.d.06.1	M00001631A:D03	77660
2363	3/24/98	342	RTA00000427F.e.12.1	M00003959C:G06	62813
2364	3/24/98	343	RTA00000527F.c.11.1	M00003817D:D12	37484
2365	3/24/98	344	RTA00000425F.p.15.1	M00001638C:H07	31680
2366	3/24/98	345	RTA00000426F.n.23.1	M00004030C:A08	18176
2367	3/24/98	346	RTA00000522F.m.19.1	M00001655C:C07	41544
2368	3/24/98	347	RTA00000522F.a.05.1	M00001567A:C04	32611
2369	3/24/98	348	RTA00000427F.i.09.1	M00004097C:E03	65916
2370	3/24/98	349	RTA00000424F.j.09.1	M00001596B:H05	74387
2371	3/24/98	350	RTA00000424F.n.11.1	M00001582C:C04	73874
2372	3/24/98	351	RTA00000523F.1.03.1	M00004927A:A02	0
2373	3/24/98	352	RTA00000527F.e.13.1	M00003826C:F05	37588
2374	3/24/98	353	RTA00000428F.a.18.1	M00005214C:A09	0
2375	3/24/98	354	RTA00000425F.j.19.1	M00001639D:G06	77925
2376	3/24/98	355	RTA00000522F.g.12.1	M00001595A:E07	78783
2377	3/24/98	356	RTA00000523F.a.07.1	M00001693A:H06	75804
2378	3/24/98	357	RTA00000425F.e.19.1	M00001629D:B10	73409
2379	3/24/98	358	RTA00000425F.n.19.1	M00001638B:C08	78324
2380	3/24/98	359	RTA00000523F.d.21.1	M00003824B:C09	33424
2381	3/24/98	360	RTA00000523F.j.03.1	M00003860A:A08	64535

Priority	SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
2382 3/24/98 361 RTA00000523F.p.09.1 M00005178A:AOR 0 2383 3/24/98 362 RTA00000523F.p.09.1 M00005178A:AO8 0 2384 3/24/98 364 RTA00000523F.m.07.1 M00005136A:D10 0 2386 3/24/98 365 RTA00000527F.k.16.1 M0000382B:B60 1015 2387 3/24/98 366 RTA00000527F.k.16.1 M0000382B:B60 1015 2388 3/24/98 367 RTA00000527F.l.19.1 M00003882B:B06 1015 2399 3/24/98 367 RTA00000524F.c.08.1 M0000148B5D:A05 41569 2390 3/24/98 370 RTA00000524F.c.08.1 M0000164B5D:A05 41569 2391 3/24/98 371 RTA00000522F.j.09.2 M00001650D:F11 78522 2393 3/24/98 372 RTA00000522F.j.19.2 M00001650D:F11 78522 2394 3/24/98 373 RTA0000052F.j.19.2 M00001650D:F11 78522 2395 3/24/98 375 RT		•				
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2410 3/24/98 389 RTA00000424F.i.15.1 M00001596A:A02 78043 2411 3/24/98 390 RTA00000524F.a.11.1 M00005210D:C09 0 2412 3/24/98 391 RTA00000522F.m.03.1 M00001654C:G09 79194 2413 3/24/98 392 RTA00000522F.a.20.1 M00001567C:E07 74070 2414 3/24/98 393 RTA00000522F.a.20.1 M00001539B:B10 74958 2415 3/24/98 394 RTA00000527F.g.14.1 M00001539B:B10 74958 2416 3/24/98 395 RTA00000527F.g.14.1 M00001578B:A02 74809 2417 3/24/98 395 RTA00000528F.g.05.2 M00001615C:E07 3770 2418 3/24/98 397 RTA00000528F.g.05.2 M00001615C:E07 3770 2419 3/24/98 398 RTA00000527F.c.16.1 M00003821A:H09 22908 2420 3/24/98 399 RTA00000525F.c.16.1 M00003821A:H09 22908 2421 3/24/98 400	2408	3/24/98				
2411 3/24/98 390 RTA00000524F.a.11.1 M00005210D:C09 0 2412 3/24/98 391 RTA00000522F.m.03.1 M00001654C:G09 79194 2413 3/24/98 392 RTA00000522F.a.20.1 M00001567C:E07 74070 2414 3/24/98 393 RTA00000522F.a.20.1 M00001539B:B10 74958 2415 3/24/98 394 RTA00000527F.g.14.1 M00001539B:B10 74958 2416 3/24/98 395 RTA00000522F.d.06.1 M00001578B:A02 74809 2417 3/24/98 396 RTA00000528F.g.05.2 M00001615C:E07 3770 2418 3/24/98 397 RTA00000527F.c.16.1 M00003874D:H07 64599 2419 3/24/98 398 RTA00000527F.c.16.1 M00003821A:H09 22908 2420 3/24/98 399 RTA00000527F.c.16.1 M00003840B:E08 63984 2422 3/24/98 401 RTA0000052F.c.16.1 M00004040A:B04 38209 2423 3/24/98 402						
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2413 3/24/98 392 RTA00000522F.a.20.1 M00001567C:E07 74070 2414 3/24/98 393 RTA00000424F.b.15.4 M00001539B:B10 74958 2415 3/24/98 394 RTA00000527F.g.14.1 M00003845D:B02 37532 2416 3/24/98 395 RTA00000522F.d.06.1 M00001578B:A02 74809 2417 3/24/98 396 RTA00000528F.g.05.2 M00001615C:E07 3770 2418 3/24/98 397 RTA00000528F.g.05.2 M00001615C:E07 3770 2419 3/24/98 398 RTA00000527F.c.16.1 M00003821A:H09 22908 2420 3/24/98 399 RTA00000527F.c.16.1 M00003840B:E08 63984 2421 3/24/98 400 RTA00000525F.c.16.1 M00003840B:E08 63984 2422 3/24/98 401 RTA00000525F.c.16.1 M000004040A:B04 38209 2423 3/24/98 402 RTA0000052F.b.18.1 M00001636A:H12 78304 2425 3/24/98 404	2411					-
2414 3/24/98 393 RTA00000424F.b.15.4 M00001539B:B10 74958 2415 3/24/98 394 RTA00000527F.g.14.1 M00003845D:B02 37532 2416 3/24/98 395 RTA00000522F.d.06.1 M00001578B:A02 74809 2417 3/24/98 396 RTA00000528F.g.05.2 M00001615C:E07 3770 2418 3/24/98 397 RTA00000528F.g.05.2 M00003974D:H07 64599 2419 3/24/98 398 RTA00000527F.c.16.1 M00003821A:H09 22908 2420 3/24/98 399 RTA00000524F.c.07.1 M00005217A:G10 0 2421 3/24/98 400 RTA00000523F.f.17.1 M00003840B:E08 63984 2422 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 2423 3/24/98 402 RTA00000527F.p.24.1 M00004031B:A06 36832 2424 3/24/98 403 RTA00000527F.b.18.1 M00001573B:A06 3460 2425 3/24/98 404	2412					
2415 3/24/98 394 RTA00000527F.g.14.1 M00003845D:B02 37532 2416 3/24/98 395 RTA00000522F.d.06.1 M00001578B:A02 74809 2417 3/24/98 396 RTA00000528F.g.05.2 M00001615C:E07 3770 2418 3/24/98 397 RTA00000427F.e.10.1 M00003874D:H07 64599 2419 3/24/98 398 RTA00000527F.c.16.1 M00003821A:H09 22908 2420 3/24/98 399 RTA00000524F.c.07.1 M00005217A:G10 0 2421 3/24/98 400 RTA00000523F.f.17.1 M00003840B:E08 63984 2422 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 2423 3/24/98 402 RTA00000527F.p.24.1 M00004031B:A06 36832 2424 3/24/98 403 RTA00000527F.b.18.1 M00001636A:H12 78304 2425 3/24/98 405 RTA00000525F.b.18.1 M00001608C:D02 75992 2427 3/24/98 406	2413					
2416 3/24/98 395 RTA00000522F.d.06.1 M00001578B:A02 74809 2417 3/24/98 396 RTA00000528F.g.05.2 M00001615C:E07 3770 2418 3/24/98 397 RTA00000427F.e.10.1 M00003974D:H07 64599 2419 3/24/98 398 RTA00000527F.c.16.1 M00003821A:H09 22908 2420 3/24/98 399 RTA00000524F.c.07.1 M00005217A:G10 0 2421 3/24/98 400 RTA00000523F.f.17.1 M00003840B:E08 63984 2422 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 2423 3/24/98 402 RTA00000527F.p.24.1 M00004031B:A06 36832 2424 3/24/98 403 RTA00000425F.n.17.1 M00001636A:H12 78304 2425 3/24/98 404 RTA00000522F.b.18.1 M00001636A:H12 75992 2426 3/24/98 405 RTA00000523F.o.07.1 M00005176A:A05 0 2428 3/24/98 406	2414					
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2418 3/24/98 397 RTA00000427F.e.10.1 M00003974D:H07 64599 2419 3/24/98 398 RTA00000527F.c.16.1 M00003821A:H09 22908 2420 3/24/98 399 RTA00000524F.c.07.1 M00005217A:G10 0 2421 3/24/98 400 RTA00000523F.f.17.1 M00003840B:E08 63984 2422 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 2423 3/24/98 402 RTA00000527F.p.24.1 M00004031B:A06 36832 2424 3/24/98 403 RTA00000425F.n.17.1 M00001636A:H12 78304 2425 3/24/98 404 RTA00000522F.b.18.1 M00001573B:A06 3460 2426 3/24/98 405 RTA00000523F.o.07.1 M00001608C:D02 75992 2427 3/24/98 406 RTA00000523F.b.08.1 M00003851B:E01 62893 2429 3/24/98 408 RTA00000522F.o.10.1 M00001638A:C08 26893 2430 3/24/98 409	2416					
2419 3/24/98 398 RTA00000527F.c.16.1 M00003821A:H09 22908 2420 3/24/98 399 RTA00000524F.c.07.1 M00005217A:G10 0 2421 3/24/98 400 RTA00000523F.f.17.1 M00003840B:E08 63984 2422 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 2423 3/24/98 402 RTA00000527F.p.24.1 M00004031B:A06 36832 2424 3/24/98 403 RTA00000425F.n.17.1 M00001636A:H12 78304 2425 3/24/98 404 RTA00000522F.b.18.1 M00001573B:A06 3460 2426 3/24/98 405 RTA00000523F.o.07.1 M00001608C:D02 75992 2427 3/24/98 406 RTA00000523F.o.07.1 M00003851B:E01 62893 2429 3/24/98 408 RTA00000522F.o.10.1 M00001638A:C08 26893 2430 3/24/98 409 RTA00000425F.i.10.1 M00001638A:C08 26893						
2420 3/24/98 399 RTA00000524F.c.07.1 M00005217A:G10 0 2421 3/24/98 400 RTA00000523F.f.17.1 M00003840B:E08 63984 2422 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 2423 3/24/98 402 RTA00000527F.p.24.1 M00004031B:A06 36832 2424 3/24/98 403 RTA00000425F.n.17.1 M00001636A:H12 78304 2425 3/24/98 404 RTA00000522F.b.18.1 M00001573B:A06 3460 2426 3/24/98 405 RTA00000425F.e.07.1 M00001608C:D02 75992 2427 3/24/98 406 RTA00000523F.o.07.1 M00005176A:A05 0 2428 3/24/98 407 RTA00000523F.b.08.1 M00003851B:E01 62893 2429 3/24/98 408 RTA00000522F.o.10.1 M00001638A:C08 26893 2430 3/24/98 409 RTA00000425F.i.10.1 M00001638A:C08 26893	2418	3/24/98				
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2425 3/24/98 404 RTA00000522F.b.18.1 M00001573B:A06 3460 2426 3/24/98 405 RTA00000425F.e.07.1 M00001608C:D02 75992 2427 3/24/98 406 RTA00000523F.o.07.1 M00005176A:A05 0 2428 3/24/98 407 RTA00000523F.h.08.1 M00003851B:E01 62893 2429 3/24/98 408 RTA00000522F.o.10.1 M00001660D:E05 78798 2430 3/24/98 409 RTA00000425F.l.10.1 M00001638A:C08 26893		3/24/98		•		
2426 3/24/98 405 RTA00000425F.e.07.1 M00001608C:D02 75992 2427 3/24/98 406 RTA00000523F.o.07.1 M00005176A:A05 0 2428 3/24/98 407 RTA00000523F.h.08.1 M00003851B:E01 62893 2429 3/24/98 408 RTA00000522F.o.10.1 M00001660D:E05 78798 2430 3/24/98 409 RTA00000425F.l.10.1 M00001638A:C08 26893						
2427 3/24/98 406 RTA00000523F.o.07.1 M00005176A:A05 0 2428 3/24/98 407 RTA00000523F.h.08.1 M00003851B:E01 62893 2429 3/24/98 408 RTA00000522F.o.10.1 M00001660D:E05 78798 2430 3/24/98 409 RTA00000425F.l.10.1 M00001638A:C08 26893						
2428 3/24/98 407 RTA00000523F.h.08.1 M00003851B:E01 62893 2429 3/24/98 408 RTA00000522F.o.10.1 M00001660D:E05 78798 2430 3/24/98 409 RTA00000425F.l.10.1 M00001638A:C08 26893						
2429 3/24/98 408 RTA00000522F.o.10.1 M00001660D:E05 78798 2430 3/24/98 409 RTA00000425F.I.10.1 M00001638A:C08 26893						
2430 3/24/98 409 RTA00000425F.I.10.1 M00001638A:C08 26893						
2431 3/24/98 410 RTA00000427F.f.16.1 M00004119D:H06 64122						
	2431	3/24/98	410	RTA00000427F.f.16.1	M00004119D:H06	64122

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
0.400	Appln	Appln	RTA00000424F.n.12.1	M00001582C:G02	41589
2432	3/24/98	411 412	RTA00000424F.ii.12.1 RTA00000425F.i.11.1	M00001362C:G02	21716
2433	3/24/98		RTA00000425F.i.11.1	M00001664B:E08	78736
2434	3/24/98	413	RTA00000423F.1.10.1 RTA00000426F.m.12.1	M00001004B:D08	63740
2435	3/24/98	414		M00004030B:D08	37746
2436	3/24/98	415	RTA00000527F.g.12.1	M00003843C.D04 M00003852B:D11	0
2437	3/24/98	416	RTA00000527F.i.12.2	M00005832B.D11 M00005213C:A01	0
2438	3/24/98	417	RTA00000524F.b.10.1	M00003213C.A01 M00001633A:G10	42255
2439	3/24/98	418	RTA00000425F.i.18.1		0
2440	3/24/98	419	RTA00000428F.b.22.1	M00005231D:B09	744 8 5
2441	3/24/98	420	RTA00000424F.j.13.1	M00001594C:H03	
2442	3/24/98	421	RTA00000523F.i.10.1	M00003855B:B09	64876
2443	3/24/98	422	RTA00000527F.f.03.1	M00003829A:B08	17788
2444	3/24/98	423	RTA00000427F.p.06.2	M00005102C:C01	0
2445	3/24/98	424	RTA00000424F.k.10.1	M00001592D:H02	73232
2446	3/24/98	425	RTA00000522F.i.07.2	M00001649A:E10	78377
2447	3/24/98	426	RTA00000424F.k.21.1	M00001614A:A04	73197
2448	3/24/98	427	RTA00000522F.b.08.1	M00001570D:E06	26915
2449	3/24/98	428	RTA00000522F.1.08.1	M00001654A:E08	78781
2450	3/24/98	429	RTA00000525F.a.14.1	M00004033B:C02	37566
2451	3/24/98	430	RTA00000424F.g.08.1	M00001482C:F09	74928
2452	3/24/98	431	RTA00000425F.1.09.1	M00001638A:B04	75251
2453	3/24/98	432	RTA00000522F.o.20.1	M00001669C:B09	74853
2454	3/24/98	433	RTA00000527F.j.04.2	M00003856A:G04	11809
2455	3/24/98	434	RTA00000522F.c.11.1	M00001576C:H02	31064
2456	3/24/98	435	RTA00000523F.c.13.1	M00003813D:B12	40668
2457	3/24/98	436	RTA00000427F.i.21.1	M00004102C:F03	65540
2458	3/24/98	437	RTA00000427F.n.10.1	M00004960B:A08	0
2459	3/24/98	438	RTA00000522F.h.02.1	M00001595C:E09	74947
2460	3/24/98	439	RTA00000522F.g.10.1	M00001595A:C07	74294
2461	3/24/98	440	RTA00000523F.o.22.1	M00005177C:B04	0
2462	3/24/98	441	RTA00000528F.g.22.2	M00001630C:F09	920
2463	3/24/98	442	RTA00000425F.d.14.1	M00001629A:H09	13417
2464	3/24/98	443	RTA00000425F.k.16.1	M00001640A:F05	75282
2465	3/24/98	444	RTA00000525F.b.09.1	M00004035B:F05	23472
2466	3/24/98	445	RTA00000522F.j.08.2	M00001650D:D10	76613
2467	3/24/98	446	RTA00000425F.f.20.1	M00001653D:H07	74071
2468	3/24/98	447	RTA00000523F.f.19.1	M00003840B:F05	34169
2469	3/24/98	448	RTA00000425F.j.18.1	M00001639D:G12	75561
2470	3/24/98	449	RTA00000426F.m.04.1	M00004028A:B10	36865
2471	3/24/98	450	RTA00000527F.g.21.1	M00003846B:C05	36028
2472	3/24/98	451	RTA00000527F.i.15.2	M00003852C:F07	14235
2473	3/24/98	452	RTA00000525F.a.22.1	M00004033D:G06	36848
2474	3/24/98	453	RTA00000522F.p.22.1	M00001671B:F02	73322
2475	3/24/98	454	RTA00000424F.d.12.2	M00001530D:E06	74342
2476	3/24/98	455	RTA00000424F.g.24.1	M00001487C:A11	79156
2477	3/24/98	456	RTA00000427F.a.10.1	M00004038B:D01	65370
2478	3/24/98	457	RTA00000426F.h.20.1	M00003905A:H11	23187
2479	3/24/98	458	RTA00000424F.d.12.3	M00001530D:E06	74342
2480	3/24/98	459	RTA00000425F.c.03.1	M00001585D:B12	74643
2481	3/24/98	460	RTA00000523F.f.16.1	M00003840B:E07	26522

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
2482	3/24/98	461	RTA00000427F.f.15.1	M00004119D:A07	66734
2483	3/24/98	462	RTA00000427F.p.13.2	M00004695B:E04	0
2484	3/24/98	463	RTA00000523F.p.16.1	M00005179D:B03	0
2485	3/24/98	464	RTA00000522F.p.18.1	M00001671A:H06	76376
2486	3/24/98	465	RTA00000528F.d.04.1	M00001570D:E07	2395
2487	3/24/98	466	RTA00000427F.d.06.1	M00003980B:C06	33446
2488	3/24/98	467	RTA00000528F.h.02.2	M00001632C:D08	1701
2489	3/24/98	468	RTA00000524F.a.18.1	M00005211A:E09	0
2490	3/24/98	469	RTA00000522F.e.20.1	M00001590B:H10	26770
2491	3/24/98	470	RTA00000427F.p.24.2	M00004897D:F03	0
2492	3/24/98	471	RTA00000528F.c.11.1	M00001486D:D12	1701
2493	3/24/98	472	RTA00000522F.g.18.1	M00001595B:H11	73226
2494	3/24/98	473	RTA00000523F.o.21.1	M00005177C:A01	0
2495	3/24/98	474	RTA00000522F.h.05.1	M00001595C:H11	73358
2496	3/24/98	475	RTA00000427F.i.06.1	M00004097B:D03	41450
2497	3/24/98	476	RTA00000425F.n.16.1	M00001636A:C02	18265
2498	3/24/98	477	RTA00000527F.1.21.1	M00003983D:H02	36439
2499	3/24/98	478	RTA00000527F.p.09.1	M00004029C:F05	7694
2500	3/24/98	479	RTA00000527F.1.23.1	M00003984A:B06	36018
2501	3/24/98	480	RTA00000424F.d.17.3	M00001455A:E11	73958
2502	3/24/98	481	RTA00000523F.j.02.1	M00003857A:H10	62853
2503	2/24/98	1132	RTA00000119A.c.12.1	M00001453A:D08	4882
2504	2/24/98	6	RTA00000119A.j.15.1	M00001460A:E11	79623
2505	2/24/98	1041	RTA00000403F.b.05.1	M00001455B:E07	74300
2506	2/24/98	994	RTA00000408F.b.04.2	M00001455A:F04	39933
2507	2/24/98	401	RTA00000132A.c.11.1	M00001454A:G03	87278
2508	2/24/98	535	RTA00000119A.g.7.1	M00001454A:F11	83580
2509	2/24/98	867	RTA00000339F.l.21.1	M00001455D:DJ1	9781
2510	2/24/98	62	RTA00000339F.n.10.1	M00001453B:F08	13719
2511	2/24/98	380	RTA00000403F.b.19.1	M00001456B:A06	22327
2512	1/28/98	288	RTA00000181AR.i.06.3	M00001452A:C07	19119
2512	2/24/98	198	RTA00000339R.I.14.1	M00001452A:C07	19119
2513	1/28/98	288	RTA00000181AR.i.06.3	M00001452A:C07	19119
2513	2/24/98	198	RTA00000339R.l.14.1	M00001452A:C07	19119
2514	2/24/98	264	RTA00000345F.j.08.1	M00001451B:A04	16731
2515	2/24/98	1125	RTA00000118A.n.5.1	M00001451A:C10	0
2516	2/24/98	229	RTA00000345F.i.09.1	M00001450A:D08	27250
2517	2/24/98	670	RTA00000131A.i.6.1	M00001450A:B08	0
2518	2/24/98	892	RTA00000339F.m.17.1	M00001453B:H12	20854
2519	2/24/98	1123	RTA00000408F.c.23.1	M00001458C:D10	42261
2520	2/24/98	680	RTA00000345F.e.11.1	M00001391C:C04	4392
2521	2/24/98	644	RTA00000119A.j.9.1	M00001460A:B12	82060
2522	2/24/98	972	RTA00000408F.d.15.1	M00001459B:C11	78467
2523	2/24/98	1081	RTA00000403F.d.02.1	M00001458D:D01	39224
2524	2/24/98	231	RTA00000408F.d.06.1	M00001458D:C11	78997
2525	2/24/98	663	RTA00000403F.b.12.1	M00001455D:A06	78775 70160
2526	2/24/98	856 743	RTA00000408F.d.02.1 RTA00000345F.i.08.1	M00001458D:A01	79169
2527	2/24/98	743 1162		M00001449D:G10	0 18247
2528	2/24/98 2/24/98	1162 841	RTA00000408F.c.10.1 RTA00000119A.i.8.1	M00001458A:A11 M00001457A:G12	18247 82593
2529	2124198	041	KIAUUUUIIYA.I.8.I	1V10000143/A:U12	04373

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	Priority	Appln			
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2531	2/24/98	750	RTA00000113A.ii.24.1 RTA00000403F.c.05.1	M00001457A:C05	74935
2532	2/24/98	750 751	RTA00000403F.c.03.1 RTA00000422F.i.02.1	M00001456C:B12	76436
		920	RTA00000422F.1.02.1 RTA00000403F.b.24.1	M00001456B:G01	78838
2533	2/24/98	920 1251	RTA00000403F.d.24.1 RTA00000408F.d.03.1	M00001458D:A02	22768
2534	2/24/98		RTA000004081.d.03.1	M00001438D:A02 M00001395A:H02	3500
2535	2/24/98	450	RTA00000118A.a.23.1 RTA00000339F.k.22.1	M00001393A:H02 M00001427C:D01	5556
2536	2/24/98	85 684	RTA00000339F.k.22.1	M00001427C:D01 M00001426D:D12	6662
2537	2/24/98		RTA00000339F.k.20.1 RTA00000118A.d.24.1	M00001426D:D12 M00001416A:H02	81488
2538	2/24/98	129	RTA00000118A.d.17.1	M00001416A:1102 M00001416A:D09	81921
2539	2/24/98	397	RTA00000118A.d.17.1 RTA00000348R.j.16.1	M00001410A:D09	7005
2540	2/24/98	158		M00001410A:D07	18
2541	2/24/98	1025	RTA00000118A.j.24.1		7568
2542	2/24/98	1005	RTA00000339F.e.17.1	M00001397D:G08	2263
2543	2/24/98	1040	RTA00000348R.o.12.1	M00001433C:F10	
2544	2/24/98	746	RTA00000345F.e.02.1	M00001395A:E03	0 38067
2545	2/24/98	517	RTA00000118A.a.2.1	M00001395A:A12	
2546	2/24/98	1065	RTA00000195R.a.06.1	M00001394A:E04	35265
2546	1/28/98	595	RTA00000195R.a.06.1	M00001394A:E04	35265
2547	1/28/98	595	RTA00000195R.a.06.1	M00001394A:E04	35265
2547	2/24/98	1065	RTA00000195R.a.06.1	M00001394A:E04	35265
2548	1/28/98	675	RTA00000179AR.b.21.3	M00001392C:D05	4366
2548	2/24/98	1264	RTA00000345F.e.13.1	M00001392C:D05	4366
2549	1/28/98	562	RTA00000196F.j.12.1	M00001396A:H03	19294
2550	2/24/98	1042	RTA00000339F.g.10.1	M00001440C:D02	6327
2551	2/24/98	706	RTA00000403F.a.09.1	M00001448B:H05	77820
2552	2/24/98	823	RTA00000119A.k.1.1	M00001460A:H11	81282
2553	2/24/98	703	RTA00000339F.n.05.1	M00001449D:B01	39648
2554	2/24/98	787	RTA00000345F.i.24.1	M00001449C:C05	0 0
2555	2/24/98	68	RTA00000339F.n.03.1	M00001449B:B03	
2556	2/24/98	440	RTA00000403F.a.18.1	M00001448D:F12 M00001448D:E12	75726 13686
2557	2/24/98	815	RTA00000403F.a.17.1		23089
2558	2/24/98	275	RTA00000353R.j.24.1	M00001428B:D01 M00001448C:E11	73952
2559	2/24/98	902	RTA00000403F.a.10.1	M00001448C.E11	5673
2560	2/24/98	1214	RTA00000339F.j.07.1	M00001428D.B10	73559
2561	2/24/98	378	RTA00000403F.a.07.1	M00001448A:E11	18808
2562	2/24/98	473	RTA00000403F.a.05.1	M00001448A:B12	23529
2563	2/24/98	128	RTA00000403F.a.04.1 RTA00000347F.c.06.1	M00001448A:B12	18846
2564	2/24/98 2/24/98	227 35	RTA00000347F.c.00.1	M00001434A:B10	5970
2565	2/24/98	442	RTA00000339F.1.13.1	M00001450A:A02	39304
2566	2/24/98	288	RTA00000347F.0.02.1	M00001430A:A02	73109
2567 2568	2/24/98	853	RTA00000403F.j.13.2	M00001448C:110	42275
2568 2569	2/24/98	249	RTA00000119A.j.10.1	M00001460A:C10	79646
2570	2/24/98	634	RTA00000119A.j.10.1	M00001487B:A11	41587
2570 2571	2/24/98	110	RTA00000418F.k.14.1	M00001487B.A11	73856
2572	2/24/98	894	RTA00000408F.k.12.1	M00001486B:D07	77246
2572 2573	2/24/98	395	RTA00000408F.j.19.2	M00001480B:D07	73752
2573 2574	2/24/98	509	RTA00000349R.g.10.1	M00001495B:B08	5777
2575	2/24/98	426	RTA00000349R.g.10.1	M00001495B:B00	74759
2576	2/24/98	101	RTA00000121A.m.2.1	M00001507A:A11	81064
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Appln	SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
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2578 2/24/98 647 RTA00000418F.b.23.1 M00001485A:C05 28767 2579 2/24/98 569 RTA00000403F.h.18.1 M00001483C:A04 39241 2581 2/24/98 707 RTA00000403F.h.11.1 M00001483B:D04 39219 2582 2/24/98 869 RTA00000403F.h.17.1 M00001483B:D04 39219 2583 2/24/98 344 RTA00000403F.h.15.2 M00001483B:H03 78935 2584 2/24/98 344 RTA00000413F.h.15.2 M00001507A:E04 17530 2585 2/24/98 1174 RTA00000121A.k.5.1 M00001507A:E05 79523 2586 2/24/98 1174 RTA00000133A.j.13.1 M00001507A:E02 16846 2587 2/24/98 184 RTA00000403F.i.23.1 M00001507A:B02 16846 2588 2/24/98 304 RTA00000403F.i.23.1 M00001507A:B02 16846 2599 2/24/98 313 RTA00000403F.i.23.1 M0000157D:B06 8811 2599 2/24/98 312	2577			RTA00000403F.i.04.1	M00001485B:D09	8930
2579 2/24/98 569 RTA00000403F.h.18.1 M00001484C:A04 39241 2580 2/24/98 236 RTA00000403F.h.12.1 M00001483B:D04 39219 2581 2/24/98 869 RTA00000403F.h.17.1 M00001483B:D04 39219 2582 2/24/98 869 RTA00000403F.h.17.2 M00001482D:H11 26856 2583 2/24/98 344 RTA00000403F.h.15.2 M00001575D:D12 26901 2584 2/24/98 768 RTA00000121A.k.5.1 M00001577A:E04 17530 2586 2/24/98 184 RTA0000013A,i.13.1 M00001507A:E04 17530 2587 2/24/98 184 RTA0000043F.h.18.2 M00001577C:B05 16846 2589 2/24/98 304 RTA0000043F.h.18.2 M00001577C:B05 16942 2590 2/24/98 313 RTA0000043F.h.23.1 M00001577C:G05 73741 2591 2/24/98 651 RTA0000043F.h.23.1 M00001576C:G05 73741 2592 2/24/98 131						
2580 2/24/98 707 RTA00000403F.h.12.1 M00001483B:D04 39219 2581 2/24/98 869 RTA00000403F.h.10.1 M00001482D:H11 26856 2583 2/24/98 869 RTA00000403F.h.17.2 M00001482D:H11 26856 2584 2/24/98 344 RTA00000403F.m.15.2 M00001507A:E04 17530 2585 2/24/98 1174 RTA00000121A.k.5.1 M00001507A:E05 79523 2586 2/24/98 1174 RTA00000133A.j.13.1 M00001507A:E05 79523 2587 2/24/98 184 RTA00000399F.j.14.1 M00001507A:E05 79523 2588 2/24/98 130 RTA00000399F.j.14.1 M00001577B:E105 16942 2589 2/24/98 304 RTA00000439F.i.08.1 M00001577D:H06 8811 2590 2/24/98 312 RTA00000439F.i.08.1 M00001576C:G05 73741 2591 2/24/98 312 RTA0000043PF.i.08.1 M00001575D:H06 8913 2593 2/24/98 312 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>						
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2586 2/24/98 1174 RTA00000121A.k.22.1 M00001507A:C05 79523 2587 2/24/98 184 RTA00000133A.j.13.1 M00001507A:B02 16846 2588 2/24/98 304 RTA00000399F.j.14.1 M00001577D:H06 8811 2590 2/24/98 304 RTA00000403F.n.18.2 M00001577D:H06 8811 2591 2/24/98 131 RTA00000418F.e.20.1 M00001576C:G05 73741 2592 2/24/98 651 RTA00000418F.e.20.1 M00001575D:B01 38927 2594 2/24/98 800 RTA0000043F.m.12.1 M00001575D:A02 16933 2595 2/24/98 800 RTA0000043F.e.08.1 M00001573D:A02 16933 2596 2/24/98 1017 RTA0000042F.e.08.1 M00001573D:A02 39020 2597 2/24/98 269 RTA0000043F.e.13.1 M00001573D:A06 39020 2597 2/24/98 101 RTA0000043F.e.13.1 M00001573C:A080 39020 2597 2/24/98 1127						
2587 2/24/98 184 RTA00000133A.j.13.1 M00001507A:B02 16846 2588 2/24/98 1230 RTA00000039F.j.14.1 M00001577D:H06 8811 2590 2/24/98 938 RTA00000403F.i.23.1 M00001487B:E10 11364 2591 2/24/98 1131 RTA00000403F.e.20.1 M00001576C:G05 73741 2592 2/24/98 312 RTA00000403F.g.11.1 M00001575D:B10 38927 2593 2/24/98 312 RTA00000403F.g.11.1 M00001575D:B10 38927 2594 2/24/98 800 RTA0000043F.g.0.1 M00001573B:G08 73442 2595 2/24/98 1017 RTA00000422F.e.08.1 M00001573B:G08 73442 2596 2/24/98 1247 RTA00000422F.e.08.1 M00001573B:G08 73442 2597 2/24/98 847 RTA0000042F.f.03.1 M00001573B:G08 73449 2598 2/24/98 847 RTA0000042F.f.03.1 M00001573B:G08 74895 2598 2/24/98 847						
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2589 2/24/98 304 RTA00000403F.n.18.2 M00001577D:H06 8811 2590 2/24/98 938 RTA00000403F.i.23.1 M00001487B:E10 11364 2591 2/24/98 651 RTA00000418F.e.20.1 M00001576C:G05 73741 2592 2/24/98 651 RTA0000043F.g.I1.1 M00001481A:H08 24238 2593 2/24/98 312 RTA000000399F.i.08.1 M00001575D:B10 38927 2594 2/24/98 800 RTA000000418F.e.03.1 M00001575D:A02 16933 2595 2/24/98 269 RTA000000418F.e.03.1 M00001573B:G08 73442 2596 2/24/98 269 RTA00000043F.i.11.1 M00001577B:F05 74895 2598 2/24/98 847 RTA000000418F.f.03.1 M00001577B:F10 78911 2600 2/24/98 910 RTA00000418F.f.03.1 M00001577B:F10 78911 2601 2/24/98 214 RTA00000120A.g.23.1 M00001465A:E10 81189 2601 2/24/98 126 </td <td></td> <td></td> <td></td> <td>•</td> <td></td> <td></td>				•		
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2611 2/24/98 71 RTA00000120A.d.24.1 M00001464A:E10 5085 2612 2/24/98 490 RTA00000120A.d.15.1 M00001464A:B02 80533 2613 2/24/98 736 RTA00000120A.c.7.1 M00001462A:D03 80985 2614 2/24/98 724 RTA00000119A.m.17.1 M00001461A:F05 79536 2615 2/24/98 1063 RTA00000132A.n.7.1 M00001466A:F08 0 2616 2/24/98 74 RTA00000408F.e.22.2 M00001476B:F08 26930 2617 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2617 1/28/98 657 RTA00000345F.d.03.1 M00001376B:A08 19230 2617 2/24/98 1137 RTA00000403F.g.06.1 M00001480C:A05 10505 2618 2/24/98 419 RTA00000408F.h.08.1 M00001480A:D03 74575 2620 2/24/98 871 RTA00000408F.b.09.1 M00001478B:H08 19700 2622 2/24/98 638						
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2613 2/24/98 736 RTA00000120A.c.7.1 M00001462A:D03 80985 2614 2/24/98 724 RTA00000119A.m.17.1 M00001461A:F05 79536 2615 2/24/98 1063 RTA00000132A.n.7.1 M00001466A:F08 0 2616 2/24/98 74 RTA00000408F.e.22.2 M00001476B:F08 26930 2617 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2617 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2617 2/24/98 1137 RTA00000345F.d.03.1 M00001376B:A08 19230 2618 2/24/98 1012 RTA00000403F.g.06.1 M00001480C:A05 10505 2619 2/24/98 419 RTA00000408F.h.08.1 M00001479C:E01 39223 2621 2/24/98 638 RTA00000418F.b.09.1 M00001478B:H08 19700 2622 2/24/98 770 RTA00000421F.f.05.1 M00001477B:E02 5266						
2614 2/24/98 724 RTA00000119A.m.17.1 M00001461A:F05 79536 2615 2/24/98 1063 RTA00000132A.n.7.1 M00001466A:F08 0 2616 2/24/98 74 RTA00000408F.e.22.2 M00001476B:F08 26930 2617 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2617 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2617 2/24/98 1137 RTA00000345F.d.03.1 M00001376B:A08 19230 2618 2/24/98 1012 RTA00000403F.g.06.1 M00001480C:A05 10505 2619 2/24/98 419 RTA00000408F.h.08.1 M00001480A:D03 74575 2620 2/24/98 871 RTA00000403F.f.23.1 M00001479C:E01 39223 2621 2/24/98 638 RTA00000418F.b.09.1 M00001478B:H08 19700 2622 2/24/98 770 RTA00000421F.f.05.1 M00001477B:E02 5266						
2615 2/24/98 1063 RTA00000132A.n.7.1 M00001466A:F08 0 2616 2/24/98 74 RTA00000408F.e.22.2 M00001476B:F08 26930 2617 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2617 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2617 2/24/98 1137 RTA00000345F.d.03.1 M00001376B:A08 19230 2618 2/24/98 1012 RTA00000403F.g.06.1 M00001480C:A05 10505 2619 2/24/98 419 RTA00000408F.h.08.1 M00001480A:D03 74575 2620 2/24/98 871 RTA00000403F.f.23.1 M00001479C:E01 39223 2621 2/24/98 638 RTA00000418F.b.09.1 M00001478B:H08 19700 2622 2/24/98 770 RTA00000421F.f.05.1 M00001477B:E02 5266						
2616 2/24/98 74 RTA00000408F.e.22.2 M00001476B:F08 26930 2617 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2617 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2617 2/24/98 1137 RTA00000345F.d.03.1 M00001376B:A08 19230 2618 2/24/98 1012 RTA00000403F.g.06.1 M00001480C:A05 10505 2619 2/24/98 419 RTA00000408F.h.08.1 M00001480A:D03 74575 2620 2/24/98 871 RTA00000403F.f.23.1 M00001479C:E01 39223 2621 2/24/98 638 RTA00000418F.b.09.1 M00001478B:H08 19700 2622 2/24/98 770 RTA00000421F.f.05.1 M00001477B:E02 5266						
2617 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2617 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2617 2/24/98 1137 RTA00000345F.d.03.1 M00001376B:A08 19230 2618 2/24/98 1012 RTA00000403F.g.06.1 M00001480C:A05 10505 2619 2/24/98 419 RTA00000408F.h.08.1 M00001480A:D03 74575 2620 2/24/98 871 RTA00000403F.f.23.1 M00001479C:E01 39223 2621 2/24/98 638 RTA00000418F.b.09.1 M00001478B:H08 19700 2622 2/24/98 770 RTA00000421F.f.05.1 M00001477B:E02 5266						
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2618 2/24/98 1012 RTA00000403F.g.06.1 M00001480C:A05 10505 2619 2/24/98 419 RTA00000408F.h.08.1 M00001480A:D03 74575 2620 2/24/98 871 RTA00000403F.f.23.1 M00001479C:E01 39223 2621 2/24/98 638 RTA00000418F.b.09.1 M00001478B:H08 19700 2622 2/24/98 770 RTA00000421F.f.05.1 M00001477B:E02 5266						
2619 2/24/98 419 RTA00000408F.h.08.1 M00001480A:D03 74575 2620 2/24/98 871 RTA00000403F.f.23.1 M00001479C:E01 39223 2621 2/24/98 638 RTA00000418F.b.09.1 M00001478B:H08 19700 2622 2/24/98 770 RTA00000421F.f.05.1 M00001477B:E02 5266						
2620 2/24/98 871 RTA00000403F.f.23.1 M00001479C:E01 39223 2621 2/24/98 638 RTA00000418F.b.09.1 M00001478B:H08 19700 2622 2/24/98 770 RTA00000421F.f.05.1 M00001477B:E02 5266				•		
2621 2/24/98 638 RTA00000418F.b.09.1 M00001478B:H08 19700 2622 2/24/98 · 770 RTA00000421F.f.05.1 M00001477B:E02 5266						
2622 2/24/98 · 770 RTA00000421F.f.05.1 M00001477B:E02 5266						
2624 2/24/98 660 RTA00000408F.e.24.2 M00001476C:C11 75002						

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
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2625	2/24/98	144	RTA00000349R.f.15.1	M00001472A:D08	75097
2626	2/24/98	774	RTA00000403F.e.24.1	M00001476B:D10	16432
2627	2/24/98	268	RTA00000403F.e.23.1	M00001476A:D11	9626
2628	2/24/98	715	RTA00000418F.b.01.1	M00001475C:G11	76040
2629	2/24/98	1073	RTA00000418F.a.10.1	M00001475B:C04	15245
2630	2/24/98	100	RTA00000339F.o.23.1	M00001473C:D09	7801
2631	2/24/98	756	RTA00000403F.g.13.1	M00001481B:D09	38718
2632	2/24/98	915	RTA00000408F.f.14.2	M00001476D:F03	73024
2633	1/28/98	389	RTA00000181AR.k.2.2	M00001453C:A11	0
2633	1/28/98	286	RTA00000181AR.k.2.3	M00001453C:A11	0
2634	1/28/98	565	RTA00000191AF.c.10.1	M00003989B:F11	40422
2635	1/28/98	449	RTA00000181AF.m.22.3	M00001455D:F09	9283
2635	1/28/98	450	RTA00000181AF.m.21.3	M00001455D:F09	9283
2636	1/28/98	449	RTA00000181AF.m.22.3	M00001455D:F09	9283
2636	1/28/98	450	RTA00000181AF.m.21.3	M00001455D:F09	9283
2637	1/28/98	449	RTA00000181AF.m.22.3	M00001455D:F09	9283
2637	1/28/98	450	RTA00000181AF.m.21.3	M00001455D:F09	9283
2638	1/28/98	449	RTA00000181AF.m.22.3	M00001455D:F09	9283
2638	1/28/98	450	RTA00000181AF.m.21.3	M00001455D:F09	9283
2639	1/28/98	390	RTA00000197AR.f.07.1	M00001457C:C11	19261
2639	1/28/98	184	RTA00000197AF.f.7.1	M00001457C:C11	19261
2640	1/28/98	598	RTA00000197F.e.10.1	M00001454B:D08	13154
2641	1/28/98	184	RTA00000197AF.f.7.1	M00001457C:C11	19261
2641	1/28/98	390	RTA00000197AR.f.07.1	M00001457C:C11	19261
2642	1/28/98	286	RTA00000181AR.k.2.3	M00001453C:A11	0
2642	1/28/98	389	RTA00000181AR.k.2.2	M00001453C:A11	0
2643	1/28/98	667	RTA00000197AF.d.16.1	M00001452A:E07	23505
2644	1/28/98	679	RTA00000197AF.d.11.1	M00001451C:E01	27260
2645	1/28/98	664	RTA00000195R.a.23.1	M00001449C:H12	86432
2646	1/28/98	594	RTA00000181AR.e.04.3	M00001448A:G09	11825
2647	1/28/98	405	RTA00000197AF.b.24.1	M00001446C:D09	23171
2648	1/28/98	572	RTA00000181AF.1.16.2	M00001454D:E05	13532
2649	1/28/98	590	RTA00000190AF.d.2.1	M00003906B:F12	2444
2650	1/28/98	675	RTA00000179AR.b.21.3	M00001392C:D05	4366
2650	2/24/98	1264	RTA00000345F.e.13.1	M00001392C:D05	4366
2651	1/28/98	486	RTA00000190AR.p.22.2	M00003979A:E11	16368
2652	1/28/98	701	RTA00000199AF.o.10.1	M00003974C:E04	0
2653	1/28/98	704	RTA00000190AF.o.12.1	M00003972D:C09	3438
2654	1/28/98	469	RTA00000190AF.n.2.1	M00003963A:E03	5650
2655	1/28/98	612	RTA00000197AR.e.22.1	M00001456A:H02	78758
2656	1/28/98	640	RTA00000190AF.f.5.1	M00003909A:H04	5015
2657	1/28/98	539	RTA00000197AR.b.13.1	M00001445B:E04	9560
2658	1/28/98	431	RTA00000199AF.k.15.1	M00003905C:G10	8275
2659	1/28/98	74 7	RTA00000190AF.c.6.1	M00003904D:D10	4780
2660	1/28/98	584	RTA00000190AR.c.03.1	M00003904C:A08	0
2660	2/24/98	1069	RTA00000346F.k.05.1	M00003904C:A08	0
2661	1/28/98	584	RTA00000190AR.c.03.1	M00003904C:A08	0
2661	2/24/98	1069	RTA00000346F.k.05.1	M00003904C:A08	0
2662	1/28/98	577	RTA00000190AF.a.24.2	M00003901B:A05	0
2663	1/28/98	639	RTA00000199AF.j.1.1	M00003881C:G09	6006

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
2664	1/28/98	649	RTA00000190AR.I.19.2	M00003946A:H10	88204
2665	1/28/98	488	RTA00000179AR.1.22.4	M00001405B:E09	4314
2665	1/28/98	481	RTA00000179AR.1.22.2	M00001405B:E09	4314
2666	1/28/98	721	RTA00000180AF.c.4.1	M00001417B:C04	5415
2667	1/28/98	744	RTA00000196F.m.4.1	M00001413A:F03	7958
2668	1/28/98	569	RTA00000196AF.I.23.1	M00001412A:E04	12052
2669	1/28/98	707	RTA00000179AF.p.15.1	M00001411D:F05	5622
2670	1/28/98	599	RTA00000179AF.o.5.1	M00001408D:D04	6172
2671	1/28/98	420	RTA00000181AF.c.11.1	M00001445D:A06	4769
2672	1/28/98	500	RTA00000179AR.m.07.5	M00001405D:D11	0
2673	1/28/98	609	RTA00000196AF.n.05.1	M00001418B:F07	12531
2673	2/24/98	1120	RTA00000353R.1.23.1	M00001418B:F07	12531
2674	1/28/98	481	RTA00000179AR.I.22.2	M00001405B:E09	4314
2674	1/28/98	488	RTA00000179AR.I.22.4	M00001405B:E09	4314
2675	1/28/98	481	RTA00000179AR.1.22.2	M00001405B:E09	4314
2675	1/28/98	488	RTA00000179AR.1.22.4	M00001405B:E09	4314
2676	1/28/98	481	RTA00000179AR.1.22.2	M00001405B:E09	4314
2676	1/28/98	488	RTA00000179AR.I.22.4	M00001405B:E09	4314
2677	1/28/98	636	RTA00000196F.k.20.1	M00001402B:F12	6324
2678	1/28/98	691	RTA00000195F.a.10.1	M00001401C:H03	6803
2679	2/24/98	161	RTA00000418F.n.22.1	M00001659D:B05	79062
2680	1/28/98	611	RTA00000196F.I.13.2	M00001408A:H04	0
2681	1/28/98	535	RTA00000196AF.n.19.1	M00001423D:D12	6881
2682	1/28/98	413	RTA00000200F.a.12.1	M00004031D:B05	16751
2683	1/28/98	580	RTA00000197F.a.12.1	M00001438B:B09	7895
2684	1/28/98	681	RTA00000180AF.1.04.2	M00001432D:F05	11159
2685	1/28/98	568	RTA00000196AF.p.01.2	M00001430A:A02	87143
2686	1/28/98	736	RTA00000196AF.o.13.1	M00001428B:A09	0
2687	1/28/98	438	RTA00000180AR.g.03.4	M00001425A:C11	9024
2687	1/28/98	95	RTA00000180AF.g.3.1	M00001425A:C11	9024
2688	1/28/98	514	RTA00000196AF.n.02.1	M00001417D:A04	39260
2689	1/28/98	741	RTA00000196AF.n.22.1	M00001424B:H04	9572
2690	1/28/98	609	RTA00000196AF.n.05.1	M00001418B:F07	12531
2690	2/24/98	1120	RTA00000353R.1.23.1	M00001418B:F07	12531
2691	1/28/98	462	RTA00000196AF.n.17.1	M00001423D:A09	12477
2692	1/28/98	477	RTA00000180AR.e.22.2	M00001423A:G05	7714
2693	1/28/98	445	RTA00000196AF.n.13.1	M00001422C:F12	8396
2694	1/28/98	696	RTA00000180AR.d.16.3	M00001419D:C10	11393
2694	2/24/98	1184	RTA00000345F.h.08.1	M00001419D:C10	11393
2695	1/28/98	696	RTA00000180AR.d.16.3	M00001419D:C10	11393
2695	2/24/98	1184	RTA00000345F.h.08.1	M00001419D:C10	11393
2696	1/28/98	541	RTA00000197AR.b.16.1	M00001445C:A08	0
2697	1/28/98	95	RTA00000180AF.g.3.1	M00001425A:C11	9024
2697	1/28/98	438	RTA00000180AR.g.03.4	M00001425A:C11	9024
2698	1/28/98	536	RTA00000193AR.a.2.3	M00004216D:D03	0
2699	1/28/98	588	RTA00000191AF.b.4.1	M00003983C:F03	14936
2700	1/28/98	401	RTA00000195F.e.04.1	M00004465B:D04	6731
2701	2/24/98	91	RTA00000355R.e.15.1	M00004316A:G09	22639
2701	1/28/98	410	RTA00000201F.a.20.1	M00004316A:G09	22639
2702	1/28/98	410	RTA00000201F.a.20.1	M00004316A:G09	22639

Priority	SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
2702 2/24/98 91 RTA00000355R.e.15.1 M00004316A:G09 22639 2703 1/28/98 716 RTA00000200F.p.05.1 M00004285C:A08 3984 2704 2/24/98 434 RTA00000200F.n.09.2 M00004249D:B08 12391 2706 2/24/98 575 RTA000001978.Fa.18.1 M00001351C:B06 5517 2707 1/28/98 472 RTA00000192AF.p.17.1 M00004216D:C03 16501 2708 1/28/98 478 RTA00000192AF.p.17.1 M00004210B:B05 7191 2710 1/28/98 753 RTA00000192AF.p.17.1 M00004210B:B05 7191 2711 1/28/98 753 RTA00000192AF.p.17.1 M0000420B:B10 5275 2711 1/28/98 730 RTA00000192AF.p.17.1 M0000420B:B10 5275 2712 1/28/98 624 RTA000000178AF.a.12.1 M000042755:B03 7866 2714 2/24/98 169 RTA000000178AF.h.2.2 M00001376B:A08 19230 2715 1/28/98 657		Priority	•			
2703 1/28/98 716 RTA00000200F.p.05.1 M00004285C:A08 3984 2704 2724/98 434 RTA00000200F.p.05.1 M00004289C:A08 3984 2705 1/28/98 528 RTA00000200F.n.09.2 M00004249D:B08 12391 2706 2/24/98 575 RTA00000345F.a.18.1 M00001351C:B06 5517 2707 1/28/98 658 RTA00000192AF.a.1.1 M00004216C:03 16501 1728/98 472 RTA00000192AF.p.17.1 M00004214C:H05 11451 2709 1/28/98 472 RTA00000192AF.o.17.1 M00004214C:H05 11451 2710 1/28/98 753 RTA00000192AF.o.17.1 M00004210B:B05 7191 2710 1/28/98 753 RTA00000192AF.o.17.1 M00004208B:F05 9061 2712 1/28/98 730 RTA00000192AF.o.15.1 M00004208B:F05 9061 2713 1/28/98 654 RTA00000192AF.o.15.1 M00004275A:B03 7866 2714 2/24/98 169 RTA00000347F.a.17.1 M00004275A:B03 7866 2714 2/24/98 656 RTA00000347F.a.17.1 M00001376B:A08 19230 2715 1/28/98 657 RTA00000178AR.h.22.3 M00001376B:A08 19230 2715 1/28/98 657 RTA0000018AR.h.22.3 M00001376B:A08 19230 2716 1/28/98 656 RTA00000178AR.h.22.2 M00001376B:A08 19230 2717 1/28/98 522 RTA00000178AR.h.17.2 M00001376A:C05 23824 2718 2/24/98 1157 RTA00000345F.c.12.1 M00001376A:C05 23824 2724/98 1155 RTA00000345F.c.12.1 M00001376A:C05 23824 2724/98 1155 RTA00000345F.c.12.1 M00001376A:C05 23824 2724/98 1155 RTA00000339F.c.04.1 M00001376B:C06 17123 2724/98 322 RTA0000039F.o.02.1 M00001376A:C05 23824 2724/98 373 RTA0000039F.o.02.1 M00001376A:C05 23824 2724/98 374 RTA0000039F.o.02.1 M00001376A:C05 23824 2724/98 373 RTA0000039F.o.02.1 M00001376A:C05 23824 2724/98 373 RTA0000039F.o.02.1 M000013602:H11 945 2724/98 373 RTA0000039F.o.02.1 M000013602:H11 945 2724/98 373 R						
2704 2/24/98 434 RTA00000348R.b.16.1 M00001347B:H04 6510 2705 1/28/98 528 RTA00000200F.n.09.2 M000004249D:B08 12391 2706 2/24/98 575 RTA00000034Fa.al.1 M00001351C:B06 5517 2707 1/28/98 658 RTA00000192AF.a.1.1 M00004210E:B05 16501 2708 1/28/98 472 RTA00000192AF.a.1.1 M00004210E:B05 1791 2710 1/28/98 478 RTA00000192AF.a.1.7.1 M00004208D:B10 5275 2711 1/28/98 563 RTA00000192AF.a.1.1.1 M00004208B:F05 9061 2712 1/28/98 624 RTA00000197AF.a.1.1 M00004208B:F05 9061 2714 2/24/98 169 RTA00000017FA.T.a.1.1 M00001376B:A08 19230 2715 1/28/98 656 RTA000000178AR.h.22.3 M00001376B:A08 19230 2715 1/28/98 657 RTA000000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 657						
2705 1/28/98 528 RTA00000200F.n.09.2 M00004249D:B08 12391 2706 2/24/98 575 RTA00000345F.a.18.1 M00001351C:B06 5517 2707 1/28/98 472 RTA00000193AF.a.1.1 M00004216D:C03 16501 2708 1/28/98 472 RTA00000192AF.p.17.1 M00004216D:C03 16501 2710 1/28/98 533 RTA00000192AF.o.16.2 M00004208D:B10 5275 2711 1/28/98 563 RTA00000192AF.o.16.2 M00004208D:B10 5275 2712 1/28/98 664 RTA00000192AF.o.16.2 M00004208D:B10 5275 2713 1/28/98 664 RTA00000192AF.o.15.1 M00004205D:F06 0 2714 2/24/98 169 RTA00000178AR.h.22.3 M00001376B:A08 19230 2715 1/28/98 657 RTA00000178AR.h.22.3 M00001376B:A08 19230 2716 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 656 <td></td> <td></td> <td></td> <td>-</td> <td></td> <td></td>				-		
2766 2/24/98 575 RTA00000345F.a.18.1 M00001351C:B06 5517 2707 1/28/98 658 RTA00000193AF.a.1.1 M00004216D:C03 16501 2708 1/28/98 472 RTA00000192AF.o.17.1 M00004210B:B05 7191 2709 1/28/98 478 RTA00000192AF.o.17.1 M0000420B:B10 5275 2710 1/28/98 553 RTA00000192AF.o.11.1 M0000420B:B10 5275 2712 1/28/98 630 RTA00000192AF.o.11.1 M00004205D:F06 0 2712 1/28/98 624 RTA00000347Fa.17.1 M00004275A:B03 7866 2714 2/24/98 169 RTA00000347Fa.17.1 M00001376B:A08 19230 2715 1/28/98 656 RTA00000347Fa.17.1 M00001376B:A08 19230 2715 1/28/98 657 RTA00000347Fa.03.1 M00001376B:A08 19230 2716 1/28/98 657 RTA00000347Fa.10.2 M00001376B:A08 19230 2716 1/28/98 652 <t< td=""><td></td><td></td><td></td><td></td><td></td><td></td></t<>						
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2711 1/28/98 563 RTA00000192AR.o.16.2 M00004208B:F05 9061 2712 1/28/98 730 RTA00000192AR.o.11.1 M00004205D:F06 0 2713 1/28/98 624 RTA00000200F.o.15.1 M000004275A:B03 7866 2714 2/24/98 169 RTA00000178AR.h.21. M00001376B:A08 19230 2715 1/28/98 656 RTA00000178AR.h.22.2 M00001376B:A08 19230 2715 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 657 RTA00000345F.d.03.1 M00001376B:A08 19230 2716 1/28/98 657 RTA00000345F.d.03.1 M00001376B:A08 19230 2716 1/28/98 522 RTA00000345F.d.12.1 M00001376A:A08 19230 2717 1/28/98 522 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 1/28/98 1095 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>						
2712 1/28/98 730 RTA00000192AF.o.11.1 M00004205D:F06 0 2713 1/28/98 624 RTA00000200F.o.15.1 M00004275A:B03 7866 2714 2/24/98 169 RTA00000347F.a.17.1 M00001376B:A08 19230 2715 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2715 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 656 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2717 1/28/98 522 RTA00000345F.c.I.2.1 M00001376B:A08 19230 2717 1/28/98 522 RTA00000345F.c.I.2.1 M00001376B:A08 19230 2718 1/28/98 522 RTA00000345F.c.I.2.1 M00001376B:A08 19230 2718 1/28/98 109	2710	1/28/98		RTA00000192AF.o.17.1		
2713 1/28/98 624 RTA00000200F.o.15.1 M00004275A:B03 7866 2714 2/24/98 169 RTA00000347F.a.17.1 M00001366D:C06 16723 2715 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2715 1/28/98 657 RTA00000345F.d.03.1 M00001376B:A08 19230 2716 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2716 1/28/98 656 RTA00000178AR.h.22.3 M00001376A:A08 19230 2717 1/28/98 522 RTA00000345F.c.12.1 M00001376A:C05 23824 2717 1/28/98 522 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 1/28/98 522 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 1/28/98 1095 RTA00000335R.h.04.1 M00001376A:C05 23824 2719 2/24/98 115				RTA00000192AR.o.16.2	t e e e e e e e e e e e e e e e e e e e	
2714 2/24/98 169 RTA00000347F.a.17.1 M00001366D:C06 16723 2715 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2715 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2715 2/24/98 1137 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 657 RTA00000178AR.h.22.3 M00001376B:A08 19230 2716 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2716 1/28/98 656 RTA00000345F.d.03.1 M00001376B:A08 19230 2717 1/28/98 522 RTA00000345F.c.12.1 M00001376A:C05 23824 2717 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 1/28/98 522 RTA00000345F.c.12.1 M00001376A:C05 23824 2719 2/24/98 1095 RTA00000358F.c.12.1 M00001376A:C05 23824 2719 2/24/98 <td< td=""><td>2712</td><td></td><td></td><td>RTA00000192AF.o.11.1</td><td>M00004205D:F06</td><td></td></td<>	2712			RTA00000192AF.o.11.1	M00004205D:F06	
2715 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2715 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2715 2/24/98 1137 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2716 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2716 1/28/98 652 RTA00000178AR.h.17.2 M00001376B:A08 19230 2717 1/28/98 522 RTA00000178AR.h.17.2 M00001376A:C05 23824 2717 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 1/28/98 522 RTA00000359F.c.02.1 M00001376A:C05 23824 2719 2/24/98 1095 RTA00000359F.c.02.1 M00001376A:C05 23824 2719 2/24/98 1155 RTA00000359F.a.02.1 M00001366D:C12 0 2721 1/28/98	2713	1/28/98				
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2716 1/28/98 657 RTA00000178AR.h.22.2 M00001376B:A08 19230 2716 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2716 2/24/98 1137 RTA00000345F.d.03.1 M00001376B:A08 19230 2717 1/28/98 522 RTA00000178AR.h.17.2 M00001376A:C05 23824 2718 1/28/98 522 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 2/24/98 1095 RTA00000353R.h.04.1 M00001376A:C05 23824 2718 2/24/98 1095 RTA00000353R.h.04.1 M00001376A:C05 23824 2719 2/24/98 1155 RTA00000353R.h.04.1 M00001376B:C06 17123 2720 1/28/98 614 RTA00000201F.f.03.1 M00001376B:C06 17123 2721 2/24/98 16 RTA00000200F.k.11.1 M00001366D:C12 0 2722 1/28/98 501 RTA00000200F.k.11.1 M00001365A:H10 3908 2724 2/24/98 322 <td>2715</td> <td>1/28/98</td> <td>657</td> <td>RTA00000178AR.h.22.2</td> <td>M00001376B:A08</td> <td></td>	2715	1/28/98	657	RTA00000178AR.h.22.2	M00001376B:A08	
2716 1/28/98 656 RTA00000178AR.h.22.3 M00001376B:A08 19230 2716 2/24/98 1137 RTA00000345F.d.03.1 M00001376B:A08 19230 2717 1/28/98 522 RTA00000178AR.h.17.2 M00001376B:A08 19230 2717 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 1/28/98 522 RTA00000345F.c.12.1 M00001376A:C05 23824 2719 2/24/98 1095 RTA00000353R.h.04.1 M00001376A:C05 23824 2719 2/24/98 1155 RTA00000353R.h.04.1 M00001375B:C06 17123 2720 1/28/98 614 RTA00000399F.a.02.1 M00001366D:C12 0 2721 2/24/98 16 RTA00000399F.a.02.1 M00001366D:C12 0 2722 1/28/98 436 RTA0000039PF.a.02.1 M00001497C:F03 9796 2723 2/24/98 114 RTA00000339F.c.05.1 M00001364B:B06 5516 2723 2/24/98 33	2715	2/24/98	1137	RTA00000345F.d.03.1	M00001376B:A08	19230
2716 2/24/98 1137 RTA00000345F.d.03.1 M00001376B:A08 19230 2717 1/28/98 522 RTA00000178AR.h.17.2 M00001376A:C05 23824 2717 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 1/28/98 522 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 2/24/98 1095 RTA00000354F.c.12.1 M00001376A:C05 23824 2719 2/24/98 1155 RTA00000354F.c.12.1 M00001376A:C05 23824 2719 2/24/98 1155 RTA00000354F.c.12.1 M00001376A:C05 23824 2720 1/28/98 614 RTA00000354F.c.12.1 M00001366D:C12 0 2721 1/28/98 436 RTA0000020R.l.1.1 M00001366D:C12 0 2722 1/28/98 501 RTA00000339F.c.05.1 M00001365A:H10 3908 2724 2/24/98 322 RTA00000339F.c.04.1 M00001362C:H11 3908 2724 1/28/98 33	2716	1/28/98	657	RTA00000178AR.h.22.2	M00001376B:A08	19230
2717 1/28/98 522 RTA00000178AR.h.17.2 M00001376A:C05 23824 2717 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 1/28/98 522 RTA000000345F.c.12.1 M00001376A:C05 23824 2718 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2719 2/24/98 1155 RTA00000353R.h.04.1 M00001376A:C05 23824 2719 2/24/98 614 RTA00000201F.f.03.1 M00001375B:C06 17123 2720 1/28/98 614 RTA00000200F.d.11.1 M00001366D:C12 0 2721 2/24/98 16 RTA00000200AF.k.11.1 M00001366D:C12 0 2722 1/28/98 501 RTA0000020AF.k.11.1 M00001365A:H10 3908 2724 1/28/98 322 RTA00000339F.c.05.1 M00001362B:H01 1805 2725 2/24/98 38 RTA00000339R.c.04.1 M00001362C:H11 945 2726 1/28/98 33	2716	1/28/98	656	RTA00000178AR.h.22.3	M00001376B:A08	19230
2717 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2718 1/28/98 522 RTA00000178AR.h.17.2 M00001376A:C05 23824 2718 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2719 2/24/98 1155 RTA00000353R.h.04.1 M00001376B:C06 17123 2720 1/28/98 614 RTA00000201F.f.03.1 M00004493B:D09 22633 2721 2/24/98 16 RTA00000399F.a.02.1 M00001366D:C12 0 2722 1/28/98 436 RTA00000200AF.k.11.1 M00004197C:F03 9796 2722 1/28/98 501 RTA000000339F.c.05.1 M00001365A:H10 3908 2724 2/24/98 322 RTA00000339F.c.04.1 M00001366B:B06 5516 2725 2/24/98 33 RTA00000339F.c.04.1 M00001362C:H11 945 2726 1/28/98 33 RTA00000339F.b.07.1 M00001362C:H11 945 2727 1/28/98 33	2716	2/24/98	1137	RTA00000345F.d.03.1	M00001376B:A08	19230
2718 1/28/98 522 RTA00000178AR.h.17.2 M00001376A:C05 23824 2718 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2719 2/24/98 1155 RTA00000353R.h.04.1 M00001375B:C06 17123 2720 1/28/98 614 RTA00000201F.f.03.1 M00004375B:C06 17123 2721 1/28/98 614 RTA00000201F.f.03.1 M00004136B:D09 22633 2721 1/28/98 436 RTA00000200AF.k.11.1 M00004197C:F03 9796 2722 1/28/98 501 RTA00000339F.c.05.1 M00001365A:H10 3908 2724 2/24/98 322 RTA00000339F.c.05.1 M00001364B:B06 5516 2725 2/24/98 88 RTA00000339F.c.04.1 M00001362D:H01 1805 2726 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33	2717	1/28/98	522	RTA00000178AR.h.17.2	M00001376A:C05	23824
2718 2/24/98 1095 RTA00000345F.c.12.1 M00001376A:C05 23824 2719 2/24/98 1155 RTA00000353R.h.04.1 M00001375B:C06 17123 2720 1/28/98 614 RTA00000201F.f.03.1 M00001493B:D09 22633 2721 2/24/98 16 RTA00000200AF.k.11.1 M00001366D:C12 0 2722 1/28/98 436 RTA00000200AF.k.11.1 M00004197C:F03 9796 2722 1/28/98 501 RTA00000200R.k.11.1 M00004197C:F03 9796 2723 2/24/98 1140 RTA00000339F.c.05.1 M00001365A:H10 3908 2724 2/24/98 322 RTA00000339F.c.04.1 M00001362C:H01 1805 2725 2/24/98 88 RTA00000339F.b.17.1 M00001362C:H11 945 2726 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33	2717	2/24/98	1095	RTA00000345F.c.12.1	M00001376A:C05	23824
2719 2/24/98 1155 RTA0000353R.h.04.1 M00001375B:C06 17123 2720 1/28/98 614 RTA00000201F.f.03.1 M00004493B:D09 22633 2721 2/24/98 16 RTA00000399F.a.02.1 M00001366D:C12 0 2722 1/28/98 436 RTA00000200AF.k.11.1 M00004197C:F03 9796 2722 1/28/98 501 RTA00000239F.c.05.1 M00001365A:H10 3908 2724 2/24/98 322 RTA00000339F.c.04.1 M00001364B:B06 5516 2725 2/24/98 888 RTA00000339R.c.04.1 M00001362D:H01 1805 2726 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000339F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000339F.b.21.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000339F.b.21.1 M00001362C:H11 945 2728 2/24/98 979 RTA00	2718	1/28/98	522	RTA00000178AR.h.17.2	M00001376A:C05	23824
2720 1/28/98 614 RTA00000201F.f.03.1 M00004493B:D09 22633 2721 2/24/98 16 RTA00000399F.a.02.1 M00001366D:C12 0 2722 1/28/98 436 RTA00000200R.k.11.1 M00004197C:F03 9796 2722 1/28/98 501 RTA00000239F.c.05.1 M00001365A:H10 3908 2723 2/24/98 322 RTA00000339F.c.04.1 M00001364B:B06 5516 2724 2/24/98 322 RTA00000339R.c.04.1 M00001362D:H01 1805 2726 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2726 2/24/98 979 RTA00000339F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000339R.b.07.1 M00001362C:H11 945 2727 2/24/98 979 RTA00000339R.b.07.1 M00001362C:H11 945 2728 2/24/98 1173 RTA000	2718	2/24/98	1095	RTA00000345F.c.12.1	M00001376A:C05	23824
2721 2/24/98 16 RTA00000399F.a.02.1 M00001366D:C12 0 2722 1/28/98 436 RTA00000200AF.k.11.1 M00004197C:F03 9796 2722 1/28/98 501 RTA00000200R.k.11.1 M00004197C:F03 9796 2723 2/24/98 1140 RTA00000339F.c.05.1 M00001365A:H10 3908 2724 2/24/98 322 RTA00000339F.c.24.1 M00001364B:B06 5516 2725 2/24/98 888 RTA00000339F.c.04.1 M00001362C:H11 945 2726 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000339F.b.07.1 M00001362C:H11 945 2727 2/24/98 979 RTA00000339F.b.07.1 M00001362C:H11 945 2728 2/24/98 973 RTA00000339F.b.22.1 M00001360A:G10 6826 2729 2/24/98 973 RTA0000	2719	2/24/98	1155	RTA00000353R.h.04.1	M00001375B:C06	17123
2722 1/28/98 436 RTA00000200AF.k.11.1 M00004197C:F03 9796 2722 1/28/98 501 RTA00000200R.k.11.1 M00004197C:F03 9796 2723 2/24/98 1140 RTA00000339F.c.05.1 M00001365A:H10 3908 2724 2/24/98 322 RTA00000339F.c.24.1 M00001362D:H01 1805 2725 2/24/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2726 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 37 RTA000000345F.b.17.1 M00001362C:H11 945 2728 2/24/98 1173 RTA000000345F.b.17.1 M00001360A:G10 6826 2729 2/24/98 973 RT	2720	1/28/98	614	RTA00000201F.f.03.1	M00004493B:D09	22633
2722 1/28/98 501 RTA00000200R.k.11.1 M00004197C:F03 9796 2723 2/24/98 1140 RTA00000339F.c.05.1 M00001365A:H10 3908 2724 2/24/98 322 RTA00000339F.c.24.1 M00001364B:B06 5516 2725 2/24/98 888 RTA00000339R.c.04.1 M00001362D:H01 1805 2726 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2726 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000345F.b.17.1 M00001362C:H11 945 2727 2/24/98 979 RTA00000339R.b.07.1 M00001362C:H11 945 2728 2/24/98 973 RTA00000339R.b.07.1 M00001360A:G10 6826 2729 2/24/98 973 RTA	2721	2/24/98	16	RTA00000399F.a.02.1	M00001366D:C12	0
2723 2/24/98 1140 RTA00000339F.c.05.1 M00001365A:H10 3908 2724 2/24/98 322 RTA00000339F.c.24.1 M00001364B:B06 5516 2725 2/24/98 888 RTA00000339F.c.04.1 M00001362D:H01 1805 2726 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2726 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 2/24/98 979 RTA00000339F.b.07.1 M00001362C:H11 945 2728 2/24/98 979 RTA00000339F.b.07.1 M00001360A:G10 6826 2729 2/24/98 973 RTA00000339F.b.22.1 M00001373D:B03 6867 2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA000000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 <td< td=""><td>2722</td><td>1/28/98</td><td>436</td><td>RTA00000200AF.k.11.1</td><td>M00004197C:F03</td><td>9796</td></td<>	2722	1/28/98	436	RTA00000200AF.k.11.1	M00004197C:F03	9796
2724 2/24/98 322 RTA00000339F.c.24.1 M00001364B:B06 5516 2725 2/24/98 888 RTA00000339R.c.04.1 M00001362D:H01 1805 2726 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2726 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2728 2/24/98 1173 RTA00000339R.b.07.1 M00001360A:G10 6826 2729 2/24/98 973 RTA00000339F.b.22.1 M00001373D:B03 6867 2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 <t< td=""><td>2722</td><td>1/28/98</td><td>501</td><td>RTA00000200R.k.11.1</td><td>M00004197C:F03</td><td>9796</td></t<>	2722	1/28/98	501	RTA00000200R.k.11.1	M00004197C:F03	9796
2725 2/24/98 888 RTA00000339R.c.04.1 M00001362D:H01 1805 2726 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2726 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2728 2/24/98 1173 RTA00000339F.b.07.1 M00001360A:G10 6826 2729 2/24/98 973 RTA00000339F.b.22.1 M00001373D:B03 6867 2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA0000020AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000020AF.g.15.1 M00004135B:G01 22898 2733 1/28/98 476 <	2723	2/24/98	1140	RTA00000339F.c.05.1	M00001365A:H10	3908
2726 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2726 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2728 2/24/98 979 RTA00000339R.b.07.1 M00001360A:G10 6826 2729 2/24/98 973 RTA00000339F.b.22.1 M00001373D:B03 6867 2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 474 RTA00000192AF.b.11.1 M00004135B:G01 22898 2734 1/28/98 735	2724	2/24/98	322	RTA00000339F.c.24.1	M00001364B:B06	5516
2726 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2727 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2728 2/24/98 1173 RTA00000339F.b.07.1 M00001360A:G10 6826 2729 2/24/98 973 RTA00000339F.b.22.1 M00001373D:B03 6867 2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200AF.g.15.1 M00004135B:G01 22898 2733 1/28/98 476 RTA0000020AF.g.15.1 M00004135B:G01 22898 2733 1/28/98 476 RTA0000020AF.g.15.1 M00004135B:G01 22898 2734 1/28/98 735	2725	2/24/98	888	RTA00000339R.c.04.1	M00001362D:H01	1805
2727 1/28/98 33 RTA00000178AR.a.20.1 M00001362C:H11 945 2727 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2728 2/24/98 1173 RTA00000339R.b.07.1 M00001360A:G10 6826 2729 2/24/98 973 RTA00000339F.b.22.1 M00001373D:B03 6867 2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 476 RTA00000192AR.d.1.3 M0000413D:H01 14507 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 752	2726	1/28/98	33	RTA00000178AR.a.20.1	M00001362C:H11	945
2727 2/24/98 979 RTA00000345F.b.17.1 M00001362C:H11 945 2728 2/24/98 1173 RTA00000339R.b.07.1 M00001360A:G10 6826 2729 2/24/98 973 RTA00000339F.b.22.1 M00001373D:B03 6867 2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 637 RTA00000200R.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 476 RTA00000192AR.d.1.3 M00004130D:H01 14507 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 752 RTA000000200R.f.10.1 M0000411D:B07 4 2736 1/28/98 516	2726	2/24/98	979	RTA00000345F.b.17.1	M00001362C:H11	945
2728 2/24/98 1173 RTA00000339R.b.07.1 M00001360A:G10 6826 2729 2/24/98 973 RTA00000339F.b.22.1 M00001373D:B03 6867 2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2731 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2732 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 474 RTA00000192AR.d.1.3 M00004135B:G01 22898 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 726 RTA00000200R.f.10.1 M00004111D:B07 4 2736 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685	2727	1/28/98	33	RTA00000178AR.a.20.1	M00001362C:H11	945
2729 2/24/98 973 RTA00000339F.b.22.1 M00001373D:B03 6867 2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2731 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2732 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 474 RTA00000192AR.d.1.3 M00004130D:H01 14507 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 726 RTA00000192AF.b.7.1 M0000411D:B07 4 2736 1/28/98 752 RTA0000020AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000355R.a.12.1 M00004159C:F09 36756 2738 2/24/98 704	2727	2/24/98	979	RTA00000345F.b.17.1	M00001362C:H11	945
2730 1/28/98 581 RTA00000191AF.p.3.2 M00004104B:F11 17 2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2731 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2732 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 474 RTA00000192AR.d.1.3 M00004130D:H01 14507 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 726 RTA00000200R.f.10.1 M0000411D:B07 4 2736 1/28/98 752 RTA00000192AF.o.7.1 M00004204D:C03 5275 2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000355R.a.12.1 M00004159C:F09 36756 2738 2/24/98 704	2728	2/24/98	1173	RTA00000339R.b.07.1	M00001360A:G10	6826
2731 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2731 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2732 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 474 RTA00000192AR.d.1.3 M00004130D:H01 14507 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 726 RTA00000200R.f.10.1 M0000411D:B07 4 2736 1/28/98 752 RTA00000192AF.o.7.1 M00004204D:C03 5275 2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000355R.a.12.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2729	2/24/98	973	RTA00000339F.b.22.1	M00001373D:B03	6867
2731 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2732 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 474 RTA00000192AR.d.1.3 M00004130D:H01 14507 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 726 RTA00000200R.f.10.1 M00004111D:B07 4 2736 1/28/98 752 RTA00000192AF.o.7.1 M00004204D:C03 5275 2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2730	1/28/98	581	RTA00000191AF.p.3.2	M00004104B:F11	17
2732 1/28/98 637 RTA00000200AF.g.15.1 M00004135B:G01 22898 2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 474 RTA00000192AR.d.1.3 M00004130D:H01 14507 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 726 RTA00000200R.f.10.1 M00004111D:B07 4 2736 1/28/98 752 RTA00000192AF.o.7.1 M00004204D:C03 5275 2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2731	1/28/98	637	RTA00000200AF.g.15.1	M00004135B:G01	22898
2732 1/28/98 476 RTA00000200R.g.15.1 M00004135B:G01 22898 2733 1/28/98 474 RTA00000192AR.d.1.3 M00004130D:H01 14507 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 726 RTA00000200R.f.10.1 M0000411D:B07 4 2736 1/28/98 752 RTA00000192AF.o.7.1 M00004204D:C03 5275 2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2731	1/28/98	476	RTA00000200R.g.15.1	M00004135B:G01	22898
2733 1/28/98 474 RTA00000192AR.d.1.3 M00004130D:H01 14507 2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 726 RTA00000200R.f.10.1 M00004111D:B07 4 2736 1/28/98 752 RTA00000192AF.o.7.1 M00004204D:C03 5275 2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2732	1/28/98	637	RTA00000200AF.g.15.1	M00004135B:G01	22898
2734 1/28/98 735 RTA00000192AF.b.11.1 M00004117A:G01 40014 2735 1/28/98 726 RTA00000200R.f.10.1 M00004111D:B07 4 2736 1/28/98 752 RTA00000192AF.o.7.1 M00004204D:C03 5275 2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2732	1/28/98	476	RTA00000200R.g.15.1	M00004135B:G01	22898
2735 1/28/98 726 RTA00000200R.f.10.1 M00004111D:B07 4 2736 1/28/98 752 RTA00000192AF.o.7.1 M00004204D:C03 5275 2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2733	1/28/98	474	RTA00000192AR.d.1.3	M00004130D:H01	14507
2736 1/28/98 752 RTA00000192AF.o.7.1 M00004204D:C03 5275 2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2734	1/28/98	735	RTA00000192AF.b.11.1	M00004117A:G01	40014
2737 1/28/98 516 RTA00000200AF.e.23.1 M00004107B:A06 14686 2738 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2735	1/28/98	726	RTA00000200R.f.10.1	M00004111D:B07	4
2738 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2736	1/28/98	752	RTA00000192AF.o.7.1	M00004204D:C03	5275
2738 2/24/98 704 RTA00000355R.a.12.1 M00004159C:F09 36756	2737	1/28/98	516	RTA00000200AF.e.23.1	M00004107B:A06	
	2738	1/28/98	685	RTA00000200F.i.9.1	M00004159C:F09	36756
2739 1/28/98 417 RTA00000200R.d.16.1 M00004085A:B02 39875	2738	2/24/98	704		M00004159C:F09	
	2739	1/28/98	417	RTA00000200R.d.16.1	M00004085A:B02	39875

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
2740	1/28/98	454	RTA00000200R.d.04.1	M00004078A:A06	5506
2741	1/28/98	551	RTA00000200AR.c.24.1	M00004076D:D04	15972
2742	1/28/98	524	RTA00000191AF.j.24.1	M00004076B:G03	0
2743	2/24/98	1166	RTA00000347F.h.01.1	M00004040A:G12	12043
2743	1/28/98	684	RTA000003171.II.0111	M00004040A:G12	12043
2744	2/24/98	1166	RTA00000347F.h.01.1	M00004040A:G12	12043
2744	1/28/98	684	RTA00000200AR.b.11.1	M00004040A:G12	12043
2745	1/28/98	415	RTA00000200AF.f.09.1	M00004111C:E11	12863
2746	1/28/98	448	RTA00000200AF.j.9.1	M00004177C:A01	8608
2747	2/24/98	446	RTA00000133A.m.19.2	M00001512A:G05	80167
2748	1/28/98	436	RTA00000133A.iii.13.2	M0000131271:003 M00004197C:F03	9796
2748	1/28/98	501	RTA000002007H.k.11.11	M00004197C:F03	9796
2749	1/28/98	436	RTA00000200AF.k.11.1	M00004197C:F03	9796
2749	1/28/98	501	RTA000002007H.k.11.1	M00004197C:F03	9796
2750	1/28/98	436	RTA00000200K.K.11.1	M00004197C:F03	9796
2750	1/28/98	501	RTA00000200AF.R.11.1	M00004197C:F03	9796
2751	1/28/98	610	RTA00000200AF.k.2.1	M00004197C:103	35924
2752	1/28/98	494	RTA00000200AF.k.1.1	M00004188D:G08	40049
2752	1/28/98	194	RTA00000200AF.R.1.1 RTA00000200R.k.01.1	M00004188C:A09	40049
2753	1/28/98	574	RTA00000200R.R.01.1 RTA00000192AF.f.3.1	M00004146C:C11	5257
2753 2754	1/28/98	604	RTA00000192AI .1.5.1 RTA00000200AF.j.15.1	M00004140C:C11	5849
2755	1/28/98	579	RTA00000200AF.j.15.1	M00004163D:E04 M00004157D:B03	22322
2756	1/28/98	634	RTA000002001.1.7.1 RTA00000192AF.j.6.1	M00004177D:B03	11494
2757	1/28/98	421	RTA00000192A1.j.0.1 RTA00000200AF.i.21.1	M00004172C:D00 M00004167D:A07	5316
2758	1/28/98	543	RTA00000200AF.i.19.1	M00004167B:A07	14722
2759	1/28/98	483	RTA00000200AI.h.19.1	M00004167A:1103	4642
2760	2/24/98	704	RTA00000152A1.31.15.11 RTA00000355R.a.12.1	M00004102C:707	36756
2760	1/28/98	685	RTA00000335K.a.12.1 RTA00000200F.i.9.1	M00004159C:F09	36756
2761	1/28/98	607	RTA00000200AF.k.12.1	M00004198B:D02	7359
2762	1/28/98	494	RTA00000200AF.k.1.1	M00004188C:A09	40049
2762	1/28/98	194	RTA00000200R.k.01.1	M00004188C:A09	40049
2763	2/24/98	554	RTA0000409F.i.03.1	M00001610A:E09	75968
2764	2/24/98	1228	RTA00000404F.h.10.1	M00001618A:A03	37148
2765	2/24/98	332	RTA00000409F.i.24.1	M00001611B:A09	76967
2766	2/24/98	1023	RTA00000404F.f.12.1	M00001611B:A05	39209
2767	2/24/98	572	RTA00000422F.I.03.1	M00001610D:D05	39147
2768	2/24/98	497	RTA00000350R.f.21.1	M00001610C:E07	22110
2769	2/24/98	557	RTA00000409F.j.05.1	M00001611C:C12	74128
2770	2/24/98	223	RTA00000404F.e.22.1	M00001610A:H05	11344
2771	2/24/98	165	RTA00000409F.j.07.1	M00001611C:H11	75190
2772	2/24/98	959	RTA00000340F.g.20.1	M00001609D:G10	4089
2773	2/24/98	737	RTA00000404F.e.15.1	M00001609B:C09	39101
2774	2/24/98	974	RTA00000340F.h.07.1	M00001608D:D11	19254
2775	2/24/98	857	RTA00000404F.e.09.1	M00001608B:A09	39121
2776	1/28/98	340	RTA00000185AF.n.8.1	M00001608B:A03	0
2776	2/24/98	75	RTA00000350R.i.22.1	M00001608B:A03	0
2777	1/28/98	340	RTA00000185AF.n.8.1	M00001608B:A03	0
2777	2/24/98	75	RTA00000350R.i.22.1	M00001608B:A03	0
2778	2/24/98	546	RTA00000409F.i.09.1	M00001610B:C07	75279
2779	2/24/98	374	RTA00000422F.m.04.1	M00001615B:A09	38702

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln	,		
2780	2/24/98	505	RTA00000121A.o.3.1	M00001511A:A02	81437
2781	2/24/98	453	RTA00000130A.h.13.1	M00001617A:A01	80790
2782	2/24/98	163	RTA00000422F.1.23.1	M00001616D:C11	4240
2783	2/24/98	889	RTA00000346F.b.16.1	M00001615C:G05	16485
2784	2/24/98	203	RTA00000404F.g.21.1	M00001615C:A11	37947
2785	2/24/98	32	RTA00000409F.j.02.1	M00001611B:E06	76417
2786	2/24/98	872	RTA00000409F.1.20.1	M00001615B:G01	74394
2787	2/24/98	978	RTA00000130A.e.20.1	M00001606A:H09	79502
2788	2/24/98	45	RTA00000409F.1.12.1	M00001615A:D06	26755
2789	2/24/98	182	RTA00000404F.g.14.1	M00001614D:B08	8858
2790	2/24/98	912	RTA00000404F.g.13.1	M00001614C:E06	9436
2791	2/24/98	1191	RTA00000340F.i.05.1	M00001614B:E08	0
2792	2/24/98	192	RTA00000421F.k.15.1	M00001613D:B03	2222
2793	2/24/98	360	RTA00000409F.j.19.1	M00001613A:F03	73792
2794	2/24/98	57	RTA00000409F.I.21.1	M00001615B:G07	73143
2795	2/24/98	354	RTA00000404F.c.03.2	M00001592C:F11	39198
2796	2/24/98	791	RTA00000399F.n.15.1	M00001594D:C03	3213
2797	2/24/98	921	RTA00000422F.j.02.1	M00001594D:B08	10368
2798	2/24/98	1114	RTA00000340F.f.22.1	M00001594B:F12	1720
2799	2/24/98	966	RTA00000340F.k.15.1	M00001594A:G09	19253
2800	2/24/98	46	RTA000004221.k.13.1 RTA00000404F.c.20.1	M00001594A:G09	39088
2801	2/24/98	955	RTA00000404F.e.06.1	M00001594A:D08	39315
2802	2/24/98	1103	RTA000004041.e.00.1 RTA00000346F.a.16.1	M00001507D:100 M00001593A:B07	12082
2802	2/24/98	540	RTA000003401.a.10.1 RTA00000418F.i.18.1	M00001595A:B07	78024
2803	2/24/98	1245	RTA000004781.1.16.1 RTA00000422F.k.22.1	M00001593C:B05	4098
2805	2/24/98	693	RTA000004221 .R.22.1 RTA00000404F.b.19.1	M00001592C:E05	39281
2805	2/24/98	1013	RTA00000404F.b.18.1	M00001592B:A04 M00001592A:H05	13669
2807	2/24/98	989	RTA000004041.0.18.1 RTA00000418F.i.12.1	M00001592A:1103 M00001592A:E02	78971
2808	2/24/98	404	RTA00000418F.1.12.11 RTA00000404F.b.11.1	M00001592A:E02 M00001591D:F06	39079
2809	2/24/98	786	RTA00000404F.b.09.1	M00001591D:C07	39166
2810	2/24/98	1147	RTA0000404F.c.18.1	M00001591D:C07	38982
2811	2/24/98	686	RTA000004041.c.18.1 RTA00000129A.k.21.1	M00001594A:C01 M00001601A:E12	82067
2812	2/24/98	1011	RTA00000129A.k.21.1 RTA00000400F.c.04.1	M00001601A:E12	6445
2812	2/24/98	702	RTA000004001.c.04.1	M00001618A:F08	82051
2814	2/24/98	425	RTA00000130A.b.5.1	M00001605A:H03	79579
2815	2/24/98	423	RTA00000130A.a.19.1	M00001605A:E09	0
2816	2/24/98	51	RTA00000130A.a.19.1 RTA00000129A.n.21.1	M00001603A:A00 M00001604A:C11	79381
2817	2/24/98	804	RTA00000129A.n.21.1 RTA00000129A.n.24.1	M00001604A:C11 M00001604A:C07	81409
2818	2/24/98	317	RTA00000129A.n.24.1 RTA00000195AF.b.21.1	M00001504A:C07 M00001595B:A09	
2818		602			39055
	1/28/98	864	RTA00000195AF.b.21.1	M00001595B:A09	39055
2819	2/24/98	317	RTA00000129A.n.17.1	M00001604A:A09	79811
2820	2/24/98 1/28/98	602	RTA00000195AF.b.21.1 RTA00000195AF.b.21.1	M00001595B:A09	39055
2820				M00001595B:A09	39055
2821	2/24/98	875 406	RTA00000129A.k.22.1	M00001601A:E02	79639
2822	2/24/98	406	RTA00000129A.k.12.1	M00001601A:A06	79322
2823	2/24/98	179	RTA00000418F.i.19.1	M00001596D:E03	79180
2824	2/24/98	759 206	RTA00000399F.o.06.1	M00001595D:G03	13574
2825	2/24/98	306	RTA00000404F.d.13.1	M00001595D:A04	39036
2826	2/24/98	1055	RTA00000346F.a.04.1	M00001607B:C05	5382
2827	2/24/98	350	RTA00000129A.p.3.1	M00001604A:B08	32644

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln	D.T		
2828	2/24/98	721	RTA00000422F.k.14.1	M00001649D:A08	0
2829	2/24/98	391	RTA00000130A.h.16.1	M00001617A:A08	80761
2830	2/24/98	962	RTA00000404F.p.05.2	M00001652D:E09	1896
2831	2/24/98	65	RTA00000404F.p.04.2	M00001652D:E05	39069
2832	2/24/98	1108	RTA00000346F.c.16.1	M00001652B:G10	9579
2833	2/24/98	1200	RTA00000422F.k.17.1	M00001652A:A01	38955
2834	2/24/98	1218	RTA00000418F.m.18.1	M00001653B:G10	76479
2835	2/24/98	171	RTA00000404F.n.20.1	M00001650A:C11	26865
2836	2/24/98	180	RTA00000400F.j.19.1	M00001653C:D10	4086
2837	2/24/98	556	RTA00000400F.i.11.1	M00001649C:H10	2587
2838	2/24/98	848	RTA00000404F.n.18.2	M00001649C:E11	37169
2839	2/24/98	29	RTA00000404F.n.16.2	M00001649C:D05	39095
2840	2/24/98	146	RTA00000404F.n.11.2	M00001649A:E11	38001
2841	2/24/98	700	RTA00000350R.m.14.1	M00001644C:B07	39171
2842	2/24/98	699	RTA00000340F.1.05.1	M00001644B:D06	38935
2843	2/24/98	479	RTA00000418F.m.10.1	M00001651A:H11	79110
2844	2/24/98	1169	RTA00000405F.b.08.1	M00001656B:E01	39182
2845	2/24/98	855	RTA00000423F.a.01.1	M00001659C:F10	39103
2846	2/24/98	363	RTA00000405F.c.11.1	M00001659A:D12	39068
2847	2/24/98	807	RTA00000418F.n.11.1	M00001658D:G12	78977
2848	2/24/98	795	RTA00000418F.n.07.1	M00001658B:A07	76316
2849	2/24/98	109	RTA00000422F.p.24.2	M00001658A:G09	5823
2850	2/24/98	696	RTA00000404F.p.12.2	M00001653B:C06	0
2851	2/24/98	369	RTA00000410F.m.05.1	M00001657B:B04	74964
2852	2/24/98	1229	RTA00000422F.n.14.1	M00001642C:G02	26787
2853	2/24/98	529	RTA00000422F.o.19.2	M00001655C:E01	13084
2854	2/24/98	327	RTA00000405F.a.11.1	M00001655A:B11	39124
2855	2/24/98	393	RTA00000418F.m.24.1	M00001654D:F12	77114
2856	2/24/98	381	RTA00000418F.m.23.1	M00001654D:F11	77195
2857	2/24/98	877	RTA00000418F.m.22.1	M00001654D:E12	74567
2858	2/24/98	166	RTA00000418F.m.19.1	M00001654D:A03	8890
2859	2/24/98	291	RTA00000405F.c.01.1	M00001657D:A04	19236
2860	2/24/98	356	RTA00000409F.m.24.1	M00001620D:H02	3942
2861	2/24/98	717	RTA00000404F.i.22.1	M00001625C:G05	39082
2862	2/24/98	648	RTA00000340F.i.13.1	M00001624B:B10	79299
2863	2/24/98	914	RTA00000138A.m.15.1	M00001624A:A03	41603
2864	2/24/98	587	RTA00000130A.o.21.1	M00001623A:F04	80218
2865	2/24/98	22	RTA00000130A.m.15.1	M00001622A:H12	81630
2866	2/24/98	767	RTA00000138A.p.10.1	M00001644A:H01	81625
2867	2/24/98	262	RTA00000409F.n.14.1	M00001621B:G05	78190
2868	2/24/98	960	RTA00000404F.I.19.2	M00001639B:H01	16196
2869	2/24/98	608	RTA00000404F.i.12.1	M00001620D:G11	39001
2870	2/24/98	342	RTA00000404F.i.02.1	M00001619D:D10	39015
2871	2/24/98	195 214	RTA00000404F.h.22.1	M00001619C:C07	18735
2872	2/24/98 2/24/98	52 52	RTA00000404F.h.19.1 RTA00000409F.m.12.1	M00001619A:E05 M00001618B:D09	8096 73490
2873		52 769	RTA00000409F.m.12.1 RTA00000340F.i.10.1	M00001618A:F10	73490 38561
2874	2/24/98	769 383	RTA00000340F.i.10.1 RTA00000404F.i.18.1	M00001618A:F10 M00001621C:H12	21912
2875	2/24/98	383 256		M00001621C:H12 M00001640A:H02	
2876	2/24/98		RTA00000404F.m.03.2		11799
2877	2/24/98	519	RTA00000404F.I.10.1	M00001638B:F10	23136

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
2878	2/24/98	646	RTA00000421F.m.14.1	M00001642A:F03	3524
2879	2/24/98	659	RTA000004217.m.14.1 RTA00000422F.m.24.1	M000016427:103	39159
2880	2/24/98	701	RTA00000422F.III.24.F	M00001641C:H07	77158
2881	2/24/98	873	RTA00000418F.I.06.1	M00001641C:F01	73317
2882	2/24/98	422	RTA00000418F.l.04.1	M00001641C:D02	74140
2883	2/24/98	766	RTA000004181.i.04.1 RTA00000404F.j.01.1	M00001641C:D02	26859
2884	2/24/98	20	RTA00000404F.m.04.2	M00001623D:G10	22720
2885	2/24/98	346	RTA000004041 in:04.2 RTA00000418F.j.08.1	M00001647X:X11	73382
2886	2/24/98	141	RTA00000418F.k.19.1	M00001629C:C11	74932
	2/24/98	373	RTA00000418F.k.19.1	M00001639C:C02	75385
2887	2/24/98	405	RTA00000418F.k.17.1	M00001639C:A10	75390
2888		63	RTA000004181.k.17.1 RTA00000404F.l.20.2	M00001639E:H05	38638
2889	2/24/98	133	RTA00000404F.1.20.2 RTA00000404F.1.20.1	M00001639B:H05	38638
2889	2/24/98			M00001639B:H05	38638
2890	2/24/98	133	RTA00000404F.1.20.1	M00001639B:H05	38638
2890	2/24/98	63	RTA00000404F.I.20.2	M00001639B:H03	0
2891	2/24/98	1261	RTA00000404F.m.17.2		73399
2892	2/24/98	626	RTA00000410F.j.01.1	M00001641B:F12	
2893	2/24/98	982	RTA00000126A.p.23.2	M00001552A:F06	80915
2894	2/24/98	196	RTA00000418F.k.10.1	M00001639A:G07	74454
2895	2/24/98	765	RTA00000137A.j.15.4	M00001559A:C08	4213
2896	2/24/98	895	RTA00000137A.j.11.4	M00001559A:A11	79752
2897	2/24/98	232	RTA00000128A.b.20.1	M00001558A:G09	79761
2898	2/24/98	152	RTA00000127A.i.20.1	M00001555A:B12	81418
2899	2/24/98	78 	RTA00000195AF.b.13.1	M00001560D:A03	12605
2899	1/28/98	59	RTA00000195AF.b.13.1	M00001560D:A03	12605
2900	2/24/98	448	RTA00000127A.a.3.1	M00001552A:H10	13232
2901	2/24/98	801	RTA00000128A.m.23.1	M00001561A:D01	81441
2902	2/24/98	499	RTA00000126A.p.18.2	M00001552A:E10	80881
2903	2/24/98	1212	RTA00000349R.o.03.1	M00001551D:H07	23006
2904	2/24/98	484	RTA00000126A.n.13.2	M00001551A:H06	79735
2905	2/24/98	240	RTA00000126A.n.7.2	M00001551A:D06	79557
2906	2/24/98	451	RTA00000126A.o.22.1	M00001551A:A11	81752
2907	2/24/98	513	RTA00000126A.k.7.2	M00001550A:E07	79866
2908	2/24/98	578	RTA00000127A.f.11.1	M00001554A:A08	81463
2909	2/24/98	372	RTA00000408F.p.24.1	M00001579A:E03	74286 74978
2910	2/24/98	985	RTA00000409F.a.08.1	M00001582D:B01 M00001582A:A03	74978 79780
2911	2/24/98	685	RTA00000129A.a.13.2	M00001582A:A03 M00001579D:H09	
2912	2/24/98	574	RTA00000403F.o.14.1		38971
2913	2/24/98	601	RTA00000403F.o.13.1	M00001579D:F04 M00001579C:H06	39049 73075
2914	2/24/98	432	RTA00000418F.g.05.1	M00001579C:H00 M00001560D:A03	12605
2915	1/28/98	59 78	RTA00000195AF.b.13.1	M00001560D:A03	12605
2915	2/24/98	7 8 491	RTA00000195AF.b.13.1 RTA00000418F.f.21.1	M00001500D:A03	75157
2916	2/24/98	612	RTA00000418F.1.21.1 RTA00000125A.k.14.1	M00001579B.F04 M00001545A:G05	79457
2917 2918	2/24/98 1/28/98	248	RTA00000123A.k.14.1 RTA00000198R.c.14.1	M00001545A:G05 M00001578D:C04	39814
	2/24/98	248 778	RTA00000198R.C.14.1 RTA00000347F.e.05.1	M00001578D:C04 M00001578D:C04	39814
2918 2919	1/28/98	248	RTA00000347F.e.03.1 RTA00000198R.c.14.1	M00001578D:C04 M00001578D:C04	39814
2919 2919	2/24/98	248 778	RTA00000198R.C.14.1 RTA00000347F.e.05.1	M00001578D:C04 M00001578D:C04	39814
2919	2/24/98	361	RTA00000347F.e.03.1 RTA00000422F.d.16.1	M00001578D:C04 M00001570C:G03	39133
2921	2/24/98	173	RTA00000422F.d.10.1	M00001570C:G05	74309
4,41	2,2,1,70				

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	Priority	Priority			
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2922	2/24/98	1195	RTA00000422F.e.23.1	M00001567D:B03	2113
2923	2/24/98	1168	RTA00000421F.b.06.1	M00001567A:B09	39037
2924	2/24/98	580	RTA00000403F.o.07.1	M00001579C:A01	7337
2925	2/24/98	531	RTA00000345F.n.12.1	M00001528A:C04	8001
2926	2/24/98	154	RTA00000340F.b.21.1	M00001533D:A08	
2927	2/24/98	19	RTA00000123A.k.23.1	M00001533A:G05	80313
2928	2/24/98	1265	RTA00000340F.d.07.1	M00001532D:A06	0
2929	2/24/98	1124	RTA00000123A.h.22.1	M00001532A:C01	17124
2930	2/24/98	1241	RTA00000408F.1.14.1	M00001530A:E10	12001
2931	2/24/98	534	RTA00000126A.g.7.1	M00001548A:H04	1902
2932	2/24/98	694	RTA00000418F.c.07.1	M00001529D:C05	73245
2933	2/24/98	1034	RTA00000124A.f.16.3	M00001536A:F11	47430
2934	2/24/98	790	RTA00000345F.n.08.1	M00001517A:B11	0
2935	2/24/98	613	RTA00000122A.j.22.1	M00001516A:F06	81151
2936	2/24/98	885	RTA00000122A.j.17.1	M00001516A:D02	62736
2937	2/24/98	1262	RTA00000122A.h.4.1	M00001514A:G03	33576
2938	2/24/98	135	RTA00000122A.d.5.1	M00001513A:F05	81155
2939	1/28/98	391	RTA00000179AF.e.20.3	M00001396A:C03	4009
2940	2/24/98	537	RTA00000408F.1.09.1	M00001530A:A09	75487
2941	2/24/98	683	RTA00000403F.j.21.1	M00001540D:E02	24723
2942	2/24/98	343	RTA00000422F.g.21.1	M00001583A:F07	17232
2943	2/24/98	226	RTA00000125A.k.10.1	M00001545A:F02	81644
2944	2/24/98	763	RTA00000135A.m.18.1	M00001545A:C03	19255
2945	2/24/98	156	RTA00000125A.k.1.1	M00001545A:B12	0
2946	2/24/98	597	RTA00000135A.I.1.2	M00001545A:B10	39426
2947	2/24/98	586	RTA00000125A.g.24.1	M00001544A:F05	80397
2948	2/24/98	467	RTA00000123A.n.13.2	M00001534A:D03	39167
2949	2/24/98	830	RTA00000347F.b.08.1	M00001541B:E05	17591
2950	2/24/98	9 97	RTA00000134A.l.9.1	M00001535A:D10	81814
2951	2/24/98	371	RTA00000403F.j.17.1	M00001539D:B10	38563
2952	2/24/98	33	RTA00000403F.j.15.1	M00001539B:G07	23840
2953	2/24/98	1209	RTA00000408F.n.05.2	M00001539A:H02	77883
2954	2/24/98	530	RTA00000408F.n.02.2	M00001539A:E01	76993
2955	2/24/98	1213	RTA00000135A.a.23.1	M00001537A:H05	27054
2956	2/24/98	347	RTA00000125A.n.4.1	M00001546A:D08	81984
2957	2/24/98	472	RTA00000135A.f.14.2	M00001542A:G12	79969
2958	2/24/98	243	RTA00000410F.c.14.1	M00001634A:H05	77809
2959	2/24/98	919	RTA00000410F.d.18.1	M00001635D:D05	75458
2960	2/24/98	825	RTA00000404F.k.22.2	M00001635D:C12	39084 39084
2960	2/24/98	364	RTA00000404F.k.22.1	M00001635D:C12	39084
2961	2/24/98	825	RTA00000404F.k.22.2	M00001635D:C12	39084
2961	2/24/98	364	RTA00000404F.k.22.1	M00001635D:C12	77561
2962	2/24/98	595	RTA00000410F.d.10.1	M00001635B:H02	76964
2963	2/24/98	175	RTA00000410F.d.09.1	M00001635B:H01	76964
2964	2/24/98	206	RTA00000410F.b.15.1	M00001633C:F09	77100
2965	2/24/98	1083	RTA00000418F.j.20.1	M00001634D:D04 M00001636A:F08	76093
2966	2/24/98	922	RTA00000410F.e.09.1		18225
2967	2/24/98	1035	RTA00000404F.k.15.1	M00001634A:B04 M00001633D:H06	77784
2968	2/24/98	1167	RTA00000410F.c.06.1	M00001633D:H06 M00001633D:G09	74099
2969	2/24/98	53	RTA00000410F.c.04.1	MINNOCCO I NUNDINI	14077

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
2970	2/24/98	567	RTA00000410F.c.02.1	M00001633D:D12	75055
2971	2/24/98	819	RTA00000410F.b.24.1	M00001633D:D09	75104
2972	2/24/98	666	RTA00000403F.o.19.1	M00001582D:F02	78615
2973	2/24/98	559	RTA00000418F.k.03.1	M00001634D:G11	78901
2974	2/24/98	999	RTA00000418F.k.04.1	M00001637A:A03	75864
2975	2/24/98	936	RTA00000121A.n.23.1	M00001511A:G01	26981
2976	2/24/98	201	RTA00000404F.I.09.1	M00001638B:E12	39176
2977	2/24/98	1160	RTA00000400F.g.02.1	M00001638B:E03	1508
2978	2/24/98	827	RTA00000410F.f.12.1	M00001637C:E03	73883
2979	2/24/98	622	RTA00000404F.1.07.1	M00001637C:C06	10798
2980	2/24/98	365	RTA00000418F.k.07.1	M00001637A:F10	75067
2981	2/24/98	248	RTA00000404F.k.24.1	M00001636A:C03	15256
2982	2/24/98	25	RTA00000418F.k.05.1	M00001637A:A06	73021
2983	2/24/98	1178	RTA00000400F.f.11.1	M00001636A:E07	4088
2984	2/24/98	1180	RTA00000404F.1.05.1	M00001636D:F09	38671
2985	2/24/98	711	RTA00000404F.l.03.2	M00001636B:G11	40272
2985	2/24/98	785	RTA00000404F.I.03.1	M00001636B:G11	40272
2986	2/24/98	785	RTA00000404F.1.03.1	M00001636B:G11	40272
2986	2/24/98	711	RTA00000404F.1.03.2	M00001636B:G11	40272
2987	2/24/98	711	RTA00000404F.1.03.2	M00001636B:G11	40272
2987	2/24/98	785	RTA00000404F.1.03.1	M00001636B:G11	40272
2988	2/24/98	711	RTA00000404F.1.03.2	M00001636B:G11	40272
2988	2/24/98	785	RTA00000404F.I.03.1	M00001636B:G11	40272
2989	2/24/98	1106	RTA00000410F.b.17.1	M00001633C:H05	77458
2990	2/24/98	253	RTA00000400F.f.18.1	M00001637A:E10	3764
2991	2/24/98	562	RTA00000401F.j.17.1	M00003901B:C05	5483
2992	2/24/98	1082	RTA00000137A.o.22.1	M00001587A:D01	0
2993	2/24/98	594	RTA00000129A.c.18.2	M00001587A:B10	37216
2994	2/24/98	891	RTA00000137A.p.12.1	M00001587A:B01	80614
2995	2/24/98	131	RTA00000418F.g.22.1	M00001585B:F01	74837
2996	2/24/98	880	RTA00000418F.g.20.1	M00001585B:C03	74626
2997	2/24/98	742	RTA00000410F.b.18.1	M00001633C:H11	76701
2998	2/24/98	879	RTA00000409F.b.19.1	M00001584D:H02	14479
2999	2/24/98	167	RTA00000399F.1.14.1	M00001590B:G08	3354
3000	2/24/98	1260	RTA00000422F.f.18.1	M00001583D:B08	24528
3000	2/24/98	1258	RTA00000403F.p.05.2	M00001583D:B08	24528
3001	2/24/98	1260	RTA00000422F.f.18.1	M00001583D:B08	24528
3001	2/24/98	1258	RTA00000403F.p.05.2	M00001583D:B08	24528
3002	2/24/98	1260	RTA00000422F.f.18.1	M00001583D:B08	24528
3002	2/24/98	1258	RTA00000403F.p.05.2	M00001583D:B08	24528
3003	2/24/98	1260	RTA00000422F.f.18.1	M00001583D:B08	24528
3003	2/24/98	1258	RTA00000403F.p.05.2	M00001583D:B08	24528
3004	2/24/98	67	RTA00000409F.a.22.1	M00001583B:F02	75200
3005	2/24/98	564	RTA00000418F.k.08.1	M00001639A:C03	18259
3006	1/28/98	282	RTA00000193AF.c.15.1	M00004248B:E08	3726
3007	2/24/98	242	RTA00000404F.j.08.1	M00001629B:B08	39066
3008	2/24/98	669	RTA00000410F.b.10.1	M00001633C:B09	74504
3009	2/24/98	725	RTA00000410F.b.07.1	M00001633C:A05	78916
3010	2/24/98	423	RTA00000410F.a.16.1	M00001633A:E06	73548
3011	2/24/98	695	RTA00000418F.j.15.1	M00001632C:H07	74855
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3012	2/24/98	901	RTA00000418F.j.14.1	M00001632C:B10	32623
3013	2/24/98	752	RTA00000410F.a.08.1	M00001632A:B10	73324
3014	2/24/98	1007	RTA00000404F.a.01.1	M00001589B:B08	19251
3015	2/24/98	1093	RTA00000340F.i.15.1	M00001629C:E07	26815
3016	2/24/98	664	RTA00000404F.a.09.1	M00001589C:E06	38985
3017	2/24/98	1246	RTA00000418F.j.11.1	M00001626C:E04	73853
3018	2/24/98	174	RTA00000404F.b.02.1	M00001591B:B12	38984
3019	2/24/98	1142	RTA00000418F.i.06.1	M00001591B:B06	75151
3020	2/24/98	740	RTA00000399F.1.19.1	M00001590D:G07	40145
3021	2/24/98	1098	RTA00000409F.d.16.1	M00001590C:F10	76090
3022	2/24/98	591	RTA00000409F.a.16.1	M00001583A:A05	73990
3023	2/24/98	1110	RTA00000404F.j.24.1	M00001631D:G05	39067
3024	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3024	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3024	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3025	1/28/98	68	RTA00000184F.k.19.1	M00001558B:D08	8022
3025	1/28/98	63	RTA00000184AF.k.19.1	M00001558B:D08	8022
3026	1/28/98	269	RTA00000183AF.k.13.1	M00001534B:C12	0
3027	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3027	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3027	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3028	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3028	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3028	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3029	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3029	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3029	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3030	1/28/98	34	RTA00000197AF.n.8.1	M00001536D:A12	4101
3031	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3031	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3031	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3032	1/28/98	106	RTA00000197AF.n.21.1	M00001540B:C09	0
3033	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3033	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3033	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09 M00001528A:F09	18957
3034	1/28/98	108	RTA00000183AR.h.23.2 RTA00000183AR.h.23.1	M00001528A:F09	18957 18957
3034	1/28/98	236 129	RTA00000183AR.ii.23.1	M00001528A:F09	18957
3034 3035	1/28/98 1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3035	1/28/98	108	RTA00000134A.d.10.1 RTA00000183AR.h.23.2	M00001528A:F09	18957
3035	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3036	1/28/98	233	RTA00000197AF.I.8.1	M00001528A:109	39954
3037	1/28/98	323	RTA00000197A1.i.o.1 RTA00000182AF.m.21.1	M00001311B:C00 M00001490C:C12	18699
3037	1/28/98	223	RTA00000192A1.iii.21.1	M00001430C:C12	0
3038	1/28/98	236	RTA000001971.1.3.1 RTA00000183AR.h.23.1	M00001488D:C10 M00001528A:F09	18957
3039	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3039	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3040	1/28/98	352	RTA00000197AF.p.3.1	M00001520A:103	7239
3041	1/28/98	301	RTA00000181AR.i.19.3	M0000135071:7105 M00001452C:B06	16970
3041	1/28/98	295	RTA00000181AR.i.19.2	M00001452C:B06	16970

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
20.42	Appln	Appln	RTA00000184F.k.19.1	M00001558B:D08	8022
3042	1/28/98	68 63	RTA00000184AF.k.19.1	M00001558B:D08	8022
3042	1/28/98	63	RTA00000184AF.k.19.1	M00001558B:D08	8022
3043	1/28/98		RTA00000184F.k.19.1	M00001558B:D08	8022
3043	1/28/98	68	RTA00000184F.k.19.1 RTA00000184F.k.12.1	M00001557D:D09	8761
3044	1/28/98	41	RTA00000184F.k.12.1 RTA00000184F.k.09.1	M00001557C:H07	7065
3045	1/28/98	150	RTA00000184F.k.09.1 RTA00000183AF.l.18.1	M00001537C:1107 M00001535D:C01	3484
3046	1/28/98	82	RTA00000184AF.i.1.1	M00001555B:C01	0
3047	1/28/98	338	RTA00000184AF.i.1.1 RTA00000182AF.i.1.3	M00001334B:C07	7033
3048	1/28/98	327	RTA00000182AF.1.1.5 RTA00000184AR.e.15.1	M00001479B.A01 M00001549C:E06	16347
3049	1/28/98	256		M00001548A:A08	4393
3050	1/28/98	99	RTA00000184AF.d.8.1	M00001546B:C05	5777
3051	1/28/98	355	RTA00000184AR.b.24.1	M00001546B:B02	39788
3052	1/28/98	322	RTA00000184AR.b.21.1	M00001541C:B07	5739
3053	1/28/98	97	RTA00000197AF.o.2.1	M00001541C:B07	0
3054	1/28/98	313	RTA00000183AF.o.11.1		7065
3055	1/28/98	42	RTA00000184F.j.21.1	M00001557A:D02	7005
3056	1/28/98	378	RTA00000181AF.k.24.3	M00001454B:C12	7005 7005
3056	1/28/98	119	RTA00000181AR.k.24.2	M00001454B:C12	7005 7005
3056	1/28/98	116	RTA00000181AR.k.24.3	M00001454B:C12	
3057	1/28/98	134	RTA00000197F.e.11.1	M00001454B:G03	2306
3057	1/28/98	298	RTA00000197AR.e.11.1	M00001454B:G03	2306
3058	1/28/98	134	RTA00000197F.e.11.1	M00001454B:G03	2306
3058	1/28/98	298	RTA00000197AR.e.11.1	M00001454B:G03	2306
3059	1/28/98	134	RTA00000197F.e.11.1	M00001454B:G03	2306
3059	1/28/98	298	RTA00000197AR.e.11.1	M00001454B:G03	2306
3060	1/28/98	116	RTA00000181AR.k.24.3	M00001454B:C12	7005 7005
3060	1/28/98	378	RTA00000181AF.k.24.3	M00001454B:C12	
3060	1/28/98	119	RTA00000181AR.k.24.2	M00001454B:C12	7005 7005
3061	1/28/98	116	RTA00000181AR.k.24.3	M00001454B:C12	7005 7005
3061	1/28/98	378	RTA00000181AF.k.24.3	M00001454B:C12	7003 7005
3061	1/28/98	119	RTA00000181AR.k.24.2	M00001454B:C12	1027
3062	1/28/98	159	RTA00000182AF.l.12.1	M00001487A:A05	7005
3063	1/28/98	119	RTA00000181AR.k.24.2	M00001454B:C12	
3063	1/28/98	116	RTA00000181AR.k.24.3	M00001454B:C12	7005 7005
3063	1/28/98	378	RTA00000181AF.k.24.3	M00001454B:C12	
3064	1/28/98	341	RTA00000181AF.1.06.2	M00001454C:C08	0 7005
3065	1/28/98	116	RTA00000181AR.k.24.3	M00001454B:C12 M00001454B:C12	7005
3065	1/28/98	119	RTA00000181AR.k.24.2	M00001454B:C12 M00001454B:C12	7005
3065	1/28/98	378	RTA00000181AF.k.24.3	M00001454B:C12	7005
3066	1/28/98	378	RTA00000181AF.k.24.3	M00001454B:C12	7005
3066	1/28/98	116	RTA00000181AR.k.24.3	M00001454B:C12	7005
3066	1/28/98	119	RTA00000181AR.k.24.2	M00001454B:C12	7005
3067	1/28/98	378	RTA00000181AF.k.24.3	M00001454B:C12	7005
3067	1/28/98	116	RTA00000181AR.k.24.3	M00001454B:C12	7005
3067	1/28/98	119	RTA00000181AR.k.24.2 RTA00000181AF.k.24.3	M00001454B:C12	7005
3068	1/28/98	378	RTA00000181AR.k.24.3	M00001454B:C12	7005
3068	1/28/98	116	RTA00000181AR.k.24.2	M00001454B:C12	7005
3068	1/28/98	119	RTA00000197AF.d.23.1	M00001454B:C12	16130
3069	1/28/98	170 491	RTA00000197AF.d.23.1 RTA00000196F.k.11.1	M00001455A.E11 M00001399C:H12	3
3070	1/28/98	471	K I MOOOOO I FOI M. I I. I	M100001377C.1112	-

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
3071	Appln 1/28/98	Appln 119	RTA00000181AR.k.24.2	M00001454D.C12	7005
3071	1/28/98	378	RTA00000181AF.k.24.2	M00001454B:C12 M00001454B:C12	7005
3071	1/28/98	116	RTA00000181AF.k.24.3	M00001454B:C12	7005
3072	1/28/98	674			7005
3072		3	RTA00000197AR.e.24.1	M00001456B:F10	39250
3072	1/28/98 2/24/98	3 78	RTA00000197AF.e.24.1	M00001456B:F10	39250
			RTA00000195AF.b.13.1	M00001560D:A03	12605
3073 3074	1/28/98 1/28/98	59	RTA00000195AF.b.13.1	M00001560D:A03	12605
3074	1/28/98	189	RTA00000197AF.h.10.1	M00001476B:F10	15554
		46	RTA00000182AF.f.13.1	M00001470C:B10	8010
3076 3077	1/28/98	200	RTA00000182AF.f.2.1	M00001469D:D02	4794
	1/28/98	325	RTA00000182AF.d.18.4	M00001467D:H05	37435
3078	1/28/98	45	RTA00000197AR.f.12.1	M00001458C:E01	3513
3079	1/28/98	298	RTA00000197AR.e.11.1	M00001454B:G03	2306
3079	1/28/98	134	RTA00000197F.e.11.1	M00001454B:G03	2306
3080	1/28/98	37	RTA00000181AF.n.15.2	M00001457A:B07	86128
3081 3082	1/28/98	7	RTA00000197AR.e.12.1	M00001454B:G07	22095
	1/28/98	674	RTA00000197AR.e.24.1	M00001456B:F10	39250
3082	1/28/98	3	RTA00000197AF.e.24.1	M00001456B:F10	39250
3083 3084	1/28/98 1/28/98	88	RTA00000197AF.e.23.1	M00001456B:C09	37157
3084 3085		243	RTA00000181AF.m.15.3	M00001455D:A11	12081
3085 3086	1/28/98	326	RTA00000197AR.e.19.1	M00001455D:A09	8047
3086	1/28/98 1/28/98	293 380	RTA00000197AF.e.13.1	M00001454C:F02	662
3088	1/28/98	206	RTA00000182AF.k.24.1	M00001485D:B10	5625
3089	1/28/98	208	RTA00000181AF.o.04.2 RTA00000187AR.h.15.2	M00001457C:C12	22205
3099	1/28/98	68	RTA0000018/AR.n.15.2 RTA00000184F.k.19.1	M00001679A:A06 M00001558B:D08	6660 8022
3090	1/28/98	63	RTA00000184AF.k.19.1	M00001558B:D08	8022 8022
3091	1/28/98	191	RTA00000184AF.k.19.1 RTA00000187AF.p.23.1	M00001338B:D08 M00003748B:F02	39804
3092	1/28/98	10	RTA00000187AF.p.25.1 RTA00000198AF.n.16.1	M00003748B:F02 M00001694C:H10	3721
3093	1/28/98	219	RTA00000198AF.m.19.1	M00001094C:1110 M00001680D:D02	40041
3093	1/28/98	32	RTA00000198R.m.19.1	M00001680D:D02	40041
3094	1/28/98	32	RTA00000198R.m.19.1	M00001680D:D02	40041
3094	1/28/98	219	RTA00000198AF.m.19.1	M00001080D:D02	40041
3095	1/28/98	317	RTA00000198AF.p.09.1	M00001000D:D02 M00003761D:E02	10473
3095	1/28/98	186	RTA00000198R.p.09.1	M00003761D:E02	10473
3096	1/28/98	219	RTA00000198AF.m.19.1	M00003701D:E02	40041
3096	1/28/98	32	RTA00000198R.m.19.1	M00001680D:D02	40041
3097	1/28/98	64	RTA00000198AF.p.12.1	M00003763D:E10	8878
3097	1/28/98	542	RTA00000198R.p.12.1	M00003763D:E10	8878
3098	1/28/98	364	RTA00000187AF.g.13.1	M00001676C:C11	2991
3099	1/28/98	430	RTA00000198R.k.23.1	M00001661B:C08	8995
3099	1/28/98	294	RTA00000198AF.k.23.1	M00001661B:C08	8995
3100	1/28/98	430	RTA00000198R.k.23.1	M00001661B:C08	8995
3100	1/28/98	294	RTA00000198AF.k.23.1	M00001661B:C08	8995
3101	1/28/98	57	RTA00000198AF.k.20.1	M00001660C:B12	22553
3102	1/28/98	368	RTA00000198AF.k.18.1	M00001660A:C12	17432
3103	1/28/98	247	RTA00000198AF.k.08.1	M00001656C:G08	17436
3104	1/28/98	219	RTA00000198AF.m.19.1	M00001680D:D02	40041
3104	1/28/98	32	RTA00000198R.m.19.1	M00001680D:D02	40041
3105	1/28/98	199	RTA00000199R.c.09.1	M00003800A:C09	16824

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
3105	1/28/98	66	RTA00000199F.c.09.2	M00003800A:C09	16824
3106	1/28/98	225	RTA00000189AF.b.5.1	M00003828A:E04	3784
3107	1/28/98	5	RTA00000195R.c.11.1	M00003811A:E03	66087
3108	1/28/98	284	RTA00000199F.d.10.2	M00003808C:B05	22049
3108	2/24/98	816	RTA00000354R.n.04.1	M00003808C:B05	22049
3109	1/28/98	284	RTA00000199F.d.10.2	M00003808C:B05	22049
3109	2/24/98	816	RTA00000354R.n.04.1	M00003808C:B05	22049
3110	1/28/98	2	RTA00000188AF.n.15.1	M00003804A:H04	0
3111	1/28/98	317	RTA00000198AF.p.09.1	M00003761D:E02	10473
3111	1/28/98	186	RTA00000198R.p.09.1	M00003761D:E02	10473
3112	1/28/98	199	RTA00000199R.c.09.1	M00003800A:C09	16824
3112	1/28/98	66	RTA00000199F.c.09.2	M00003800A:C09	16824
3113	1/28/98	487	RTA00000198F.i.8.1	M00001639A:F10	9807
3113	1/28/98	277	RTA00000198AR.i.08.1	M00001639A:F10	9807
3114	1/28/98	66	RTA00000199F.c.09.2	M00003800A:C09	16824
3114	1/28/98	199	RTA00000199R.c.09.1	M00003800A:C09	16824
3115	1/28/98	224	RTA00000188AF.m.11.1	M00003799A:D09	0
3116	1/28/98	58	RTA00000199F.b.01.2	M00003778A:D08	19118
3117	1/28/98	216	RTA00000188AF.g.9.1	M00003774B:B08	4959
3118	1/28/98	201	RTA00000198AF.p.18.1	M00003769B:D03	23081
3119	1/28/98	542	RTA00000198R.p.12.1	M00003763D:E10	8878
3119	1/28/98	64	RTA00000198AF.p.12.1	M00003763D:E10	8878
3120	1/28/98	199	RTA00000199R.c.09.1	M00003800A:C09	16824
3120	1/28/98	66	RTA00000199F.c.09.2	M00003800A:C09	16824
3121	1/28/98	146	RTA00000185AF.a.19.2	M00001571C:H06	5749
3122	1/28/98	248	RTA00000198R.c.14.1	M00001578D:C04	39814
3122	2/24/98	778	RTA00000347F.e.05.1	M00001578D:C04	39814
3123	1/28/98	248	RTA00000198R.c.14.1	M00001578D:C04	39814
3123	2/24/98	778	RTA00000347F.e.05.1	M00001578D:C04	39814
3124	1/28/98	147	RTA00000185AF.c.24.2	M00001578B:E04	23001
3125	1/28/98	195	RTA00000198AF.c.10.1	M00001577B:H02	77149
3126	1/28/98	171	RTA00000198R.c.07.1	M00001575D:G05	19181
3126	1/28/98	525	RTA00000198AF.c.7.1	M00001575D:G05	19181
3127	1/28/98	172	RTA00000186AF.p.09.2	M00001655C:E04	6879
3128	1/28/98	230	RTA00000185AR.b.18.1	M00001575B:C09	12171
3129	1/28/98	192	RTA00000185AF.m.7.1	M00001605C:D12	39804
3130	1/28/98	19	RTA00000185AF.a.8.1	M00001570D:A03	4868
3131	1/28/98	492	RTA00000198AF.b.8.1	M00001567C:H12	22636
3131	1/28/98	23	RTA00000198R.b.08.1	M00001567C:H12	22636
3132	1/28/98	23	RTA00000198R.b.08.1	M00001567C:H12	22636
3132	1/28/98	492	RTA00000198AF.b.8.1	M00001567C:H12	22636
3133	1/28/98	357	RTA00000184AF.o.15.1	M00001564D:C09	0
3134	1/28/98	30	RTA00000184AR.n.07.2	M00001561C:F06	0
3135	1/28/98	59	RTA00000195AF.b.13.1	M00001560D:A03	12605
3135	2/24/98	78	RTA00000195AF.b.13.1	M00001560D:A03	12605
3136	1/28/98	525	RTA00000198AF.c.7.1	M00001575D:G05	19181
3136	1/28/98	171	RTA00000198R.c.07.1	M00001575D:G05	19181
3137	1/28/98	303	RTA00000186AR.e.03.3	M00001623D:C10	22110
3138	1/28/98	295	RTA00000181AR.i.19.2	M00001452C:B06	16970
3138	1/28/98	301	RTA00000181AR.i.19.3	M00001452C:B06	16970

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
3139	Appln 1/28/98	Appln 232	RTA00000186AF.j.03.2	M00001638A:E07	0
3139	1/28/98	309	RTA00000188AF.h.12.1	M00001638A:E07	9503
	1/28/98	268	RTA00000198AF.h.01.2	M00001632A:F12	0
3141 3142	1/28/98	267	RTA00000186AF.g.11.2	M00001632A:112 M00001630B:H09	5214
3142	1/28/98	83	RTA00000186AF.f.24.2	M00001630B:H09	0
3143	1/28/98	336	RTA00000186AF.f.24.1	M00001629B:E06	0
3143	1/28/98	222	RTA00000185AF.i.4.1	M00001594A:B12	13942
3145	1/28/98	217	RTA00000198AF.h.3.1	M00001625D:C07	22562
3146	1/28/98	196	RTA00000198F.e.10.1	M00001529B:E09	9727
3147	1/28/98	372	RTA000001361.c.10.1	M00001533B:G07	22187
3148	1/28/98	302	RTA00000186AF.d.1.2	M00001623D:G07	40044
3149	1/28/98	262	RTA00000186AF.c.17.1	M00001621C:C00	8551
3150	1/28/98	358	RTA00000198AF.g.7.1	M00001616C:C09	13386
3151	1/28/98	166	RTA00000198AF.f.21.1	M00001614D:D09	22676
3151	1/28/98	277	RTA00000198AR.i.08.1	M00001614D:509	9807
3152	1/28/98	487	RTA00000198F.i.8.1	M00001639A:F10	9807
3153	1/28/98	336	RTA00000186AF.f.24.1	M00001629B:E06	0
3153	1/28/98	83	RTA00000186AF.f.24.2	M00001629B:E06	Ŏ
3154	1/28/98	352	RTA00000197AF.p.3.1	M00001550A:A03	7239
3155	1/28/98	251	RTA00000192AF.n.13.1	M00004197D:H01	8210
3156	1/28/98	41	RTA00000184F.k.12.1	M00001557D:D09	8761
3157	1/28/98	731	RTA00000184F.k.02.1	M00001557B:H10	5192
3158	1/28/98	42	RTA00000184F.j.21.1	M00001557A:D02	7065
3159	1/28/98	42	RTA00000184F.j.21.1	M00001557A:D02	7065
3160	1/28/98	302	RTA00000186AF.d.1.2	M00001621C:C08	40044
3161	1/28/98	560	RTA00000184AF.i.23.3	M00001556A:F11	1577
3162	1/28/98	558	RTA00000186AR.h.14.1	M00001632D:H07	0
3163	1/28/98	256	RTA00000184AR.e.15.1	M00001549C:E06	16347
3164	1/28/98	682	RTA00000125A.j.16.1	M00001544A:E06	0
3165	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3165	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3165	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3166	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3166	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3166	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3167	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3167	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3167	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3168	2/24/98	531	RTA00000345F.n.12.1	M00001528A:C04	7337
3169	1/28/98	324	RTA00000184F.j.06.1	M00001556B:G02	11294
3170	2/24/98	604	RTA00000351R.c.13.1	M00003747D:C05	11476
3171	1/28/98	301	RTA00000181AR.i.19.3	M00001452C:B06	16970
3171	1/28/98	295	RTA00000181AR.i.19.2	M00001452C:B06	16970
3172	1/28/98	231	RTA00000192AF.1.13.2	M00004185C:C03	11443
3173	1/28/98	634	RTA00000192AF.j.6.1	M00004172C:D08	11494
3174	1/28/98	165	RTA00000192AF.g.23.1	M00004157C:A09	6455
3175	1/28/98	574	RTA00000192AF.f.3.1	M00004146C:C11	5257
3176	1/28/98	146	RTA00000185AF.a.19.2	M00001571C:H06	5749
3177	1/28/98	651	RTA00000189AR.d.22.2	M00003844C:B11	6539
3178	1/28/98	161	RTA00000183AF.e.23.2	M00001506D:A09	0

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			4 0
3179	1/28/98	475	RTA00000187AR.m.3.3	M00001682C:B12	17055
3180	2/24/98	39	RTA00000187AF.l.7.1	M00001680D:F08	10539
3181	1/28/98	228	RTA00000187AR.h.15.2	M00001679A:A06	6660
3182	2/24/98	465	RTA00000350R.p.18.1	M00001676B:F05	11460
3183	1/28/98	575	RTA00000186AF.1.12.2	M00001645A:C12	19267
3184	2/24/98	700	RTA00000350R.m.14.1	M00001644C:B07	39171
3185	1/28/98	261	RTA00000192AF.a.24.1	M00004114C:F11	13183
3186	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3186	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3186	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3187	1/28/98	398	RTA00000177AR.l.13.3	M00001353A:G12	8078
3188	1/28/98	645	RTA00000177AF.k.9.1	M00001352A:E02	16245
3189	1/28/98	283	RTA00000177AF.i.8.4	M00001350A:H01	7187
3190	1/28/98	361	RTA00000177AR.g.16.4	M00001347A:B10	13576
3191	1/28/98	680	RTA00000177AF.f.10.1	M00001345A:E01	6420
3192	1/28/98	632	RTA00000183AR.g.03.1	M00001512D:G09	3956
3192	1/28/98	630	RTA00000183AR.g.03.2	M00001512D:G09	3956
3193	1/28/98	702	RTA00000177AR.b.8.5	M00001340B:A06	17062
3194	1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3194	1/28/98	493	RTA00000177AF.m.17.1	M00001355B:G10	14391
3194	1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3195	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3195	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3195	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3196	1/28/98	129	RTA00000134A.d.10.1	M00001528A:F09	18957
3196	1/28/98	236	RTA00000183AR.h.23.1	M00001528A:F09	18957
3196	1/28/98	108	RTA00000183AR.h.23.2	M00001528A:F09	18957
3197	1/28/98	435	RTA00000182AR.c.22.1	M00001467A:D08	16283
3198	1/28/98	635	RTA00000181AF.p.7.3	M00001460A:E01	38773
3199	1/28/98	362	RTA00000197AR.c.24.1	M00001450A:B12	82498
3200	2/24/98	442	RTA00000347F.b.02.1	M00001450A:A02	39304
3201	1/28/98	265	RTA00000177AF.e.14.1	M00001343D:H07	23255
3202	1/28/98	270	RTA00000178R.1.08.1	M00001383A:C03	39648
3203	1/28/98	472	RTA00000192AF.p.17.1	M00004214C:H05	11451
3204	1/28/98	603	RTA00000183AR.d.11.3	M00001504D:G06	6420
3205	1/28/98	519	RTA00000183AF.a.24.2	M00001499B:A11	10539
3206	1/28/98	435	RTA00000182AR.c.22.1	M00001467A:D08	16283
3207	2/24/98	158	RTA00000348R.j.16.1	M00001410A:D07	7005
3208	1/28/98	411	RTA00000179AF.j.13.3	M00001400B:H06	0
3209	1/28/98	742	RTA00000177AF.m.1.1	M00001353D:D10	14929
3210	1/28/98	270	RTA00000178R.l.08.1	M00001383A:C03	39648
3211	1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3211	1/28/98	493	RTA00000177AF.m.17.1	M00001355B:G10	14391
3211	1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3212	1/28/98	297	RTA00000178AF.f.9.3	M00001371C:E09	7172
3213	1/28/98	33	RTA00000178AR.a.20.1	M00001362C:H11	945
3213	2/24/98	979	RTA00000345F.b.17.1	M00001362C:H11	945
3214	1/28/98	33	RTA00000178AR.a.20.1	M00001362C:H11	945
3214	2/24/98	979	RTA00000345F.b.17.1	M00001362C:H11	945
3215	1/28/98	466	RTA00000177AF.p.20.1	M00001361A:A05	4141
			201		

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
2216	Appln	Appln 330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3216	1/28/98 1/28/98	493	RTA00000177AR.m.17.4 RTA00000177AF.m.17.1	M00001355B:G10	14391
3216	1/28/98	337	RTA00000177AR.m.17.1 RTA00000177AR.m.17.3	M00001355B:G10	14391
3216	1/28/98	632	RTA00000177AR.iii.17.5 RTA00000183AR.g.03.1	M00001533D:G10	3956
3217 3217	1/28/98	630	RTA00000183AR.g.03.2	M00001512D:G09	3956
	1/28/98	391	RTA00000179AF.e.20.3	M00001312B:G07	4009
3218	1/28/98	370	RTA00000179AF.c.15.3	M00001390A:C05	2995
3219 3219	1/28/98	460	RTA00000179AF.c.15.1	M00001392D:H06	2995
3219	1/28/98	47	RTA00000177AF.c.13.1 RTA00000192AF.m.12.1	M00001392D:H00	0
3220	1/28/98	438	RTA00000192A1.iii.12.11 RTA00000180AR.g.03.4	M00001131B:B11	9024
3221	1/28/98	95	RTA00000180AF.g.3.1	M00001425A:C11	9024
3222	1/28/98	438	RTA00000180AR.g.03.4	M00001425A:C11	9024
3222	1/28/98	438 95	RTA00000180AR.g.3.1	M00001425A:C11	9024
	1/28/98	320	RTA00000196AF.m.13.1	M00001425A:E11	16290
3223	1/28/98	365	RTA00000196F.1.20.2	M00001419B:G05	22678
3224		303 80	RTA00000196AF.p.13.2	M00001410B:G05 M00001432A:E06	6125
3225	1/28/98	80 179	RTA00000179AF.f.20.3	M00001432A.E00 M00001397B:B09	16154
3226	1/28/98	379	RTA00000179AF.1.20.3 RTA00000180AF.1.06.2	M00001397B:B09 M00001433A:G07	5625
3227	1/28/98 1/28/98	379 460	RTA00000179AF.c.15.1	M00001433A:G07 M00001392D:H06	2995
3228		370	RTA00000179AF.c.15.1 RTA00000179AF.c.15.3	M00001392D:H06	2995
3228	1/28/98	370 107	RTA00000179AF.C.13.3 RTA00000196R.i.13.1	M00001392D:1100 M00001390A:A09	9857
3229	1/28/98	107	RTA00000178AR.m.19.5	M00001390A:A09	0
3230	1/28/98	377	RTA00000178AF.m.19.1	M00001384D:H07	0
3230	1/28/98	120	RTA00000178AR.m.19.5	M00001384D:H07	0
3231	1/28/98	377	RTA00000178AF.m.19.1	M00001384D:H07	0
3231	1/28/98 1/28/98	384	RTA00000178AF.h.17.1	M00001384D:F107	39215
3232		384 182	RTA00000196AF.h.16.1	M00001384C:F03	39895
3233 3234	1/28/98 1/28/98	105	RTA00000179AF.g.12.3	M00001384C.L03	36390
3234	1/28/98	252	RTA00000177AI .g.12.3	M0000133671:Q03	8
3235	1/28/98	253	RTA00000181AF.e.17.3	M00001448D:C09	8
3236	1/28/98	301	RTA00000181AR.i.19.3	M000014102:009	16970
3236	1/28/98	295	RTA00000181AR.i.19.2	M00001452C:B06	16970
3237	1/28/98	288	RTA00000181AR.i.06.3	M00001452A:C07	19119
3237	2/24/98	198	RTA00000339R.l.14.1	M00001452A:C07	19119
3238	1/28/98	288	RTA00000181AR.i.06.3	M00001452A:C07	19119
3238	2/24/98	198	RTA00000339R.I.14.1	M00001452A:C07	19119
3239	1/28/98	109	RTA00000197AF.d.12.1	M00001451D:C10	39546
3240	1/28/98	149	RTA00000181AR.h.06.3	M00001450D:D04	87226
3241	1/28/98	75	RTA00000180AR.h.19.2	M00001428A:H10	84182
3242	1/28/98	21	RTA00000131A.g.19.2	M00001449A:G10	36535
3243	1/28/98	308	RTA00000178AF.j.20.1	M00001380C:E05	15066
3244	1/28/98	253	RTA00000181AF.e.17.3	M00001448D:C09	8
3244	1/28/98	252	RTA00000181AF.e.18.3	M00001448D:C09	8 ,
3245	1/28/98	252	RTA00000181AF.e.18.3	M00001448D:C09	8
3245	1/28/98	253	RTA00000181AF.e.17.3	M00001448D:C09	8
3246	1/28/98	253	RTA00000181AF.e.17.3	M00001448D:C09	8
3246	1/28/98	252	RTA00000181AF.e.18.3	M00001448D:C09	8
3247	1/28/98	136	RTA00000197AF.c.10.1	M00001448B:F06	10400
3248	1/28/98	177	RTA00000197AF.c.3.1	M00001447C:C01	3145
3249	1/28/98	204	RTA00000180AR.o.5.2	M00001437D:C04	7848

Applin Applin 1/28/98 362 RTA00000197AR.c.24.1 M00001450A:B12 82498 3251 1/28/98 342 RTA00000196AF.b.15.1 M00001334C:B06 21256 3252 1/28/98 342 RTA00000196AF.d.10.1 M00001334C:B06 21256 3253 1/28/98 113 RTA00000196AF.d.09.1 M00001354C:B06 21256 3254 1/28/98 463 RTA00000177AR.k.23.4 M00001352D:D02 35550 3255 1/28/98 168 RTA00000177AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 168 RTA00000177AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 168 RTA00000177AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 135 RTA00000177AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 336 RTA00000177AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 337 RTA00000177AR.m.17.1 M000013352D:D02 35550 3255 1/28/98 337 RTA00000177AF.m.17.1 M00001338A:D04 12193 3259 1/28/98 330 RTA00000177AR.m.17.3 M00001355B:G10 14391 3259 1/28/98 330 RTA00000177AR.m.17.3 M00001355B:G10 14391 3260 1/28/98 361 RTA00000177AR.m.17.3 M00001355B:G10 14391 3266 1/28/98 361 RTA00000177AR.e.14.1 M00001347A:B10 31576 3266 1/28/98 56 RTA00000177AF.e.14.1 M00001347A:B10 31576 3266 1/28/98 48 RTA00000177AF.e.9.1 M00001343D:C04 37442 3266 1/28/98 38 RTA00000177AF.e.9.1 M00001350D:H01 31576 3266 1/28/98 38 RTA00000177AF.e.9.1 M00001350D:H01 31576 3266 1/28/98 38 RTA00000177AF.e.8.3 M00001350D:H01 31576 3267 1/28/98 38 RTA00000177AF.e.8.3 M00001350D:H01 31576 3266 1/28/98 38 RTA00000177AF.e.8.3 M00001350D:H01 31576 3266 1/28/98 38 RTA00000177AF.e.8.3 M00001350D:H01 31876 3266 1/28/98 38 RTA00000177AF.e.8.3 M00001350D:H01 3188 3268 1/28/98 337 RTA00000177AF.e.8.1 M00001350D:H01 32774	SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
3251 1/28/98 81 RTA00000196AF.b.15.1 M00001347B:E01 5102 3252 1/28/98 342 RTA00000196AF.d.10.1 M000001354B:B10 16934 3253 1/28/98 113 RTA00000196AF.d.10.1 M000001354B:B10 16934 3254 1/28/98 463 RTA00000177AR.k.23.4 M00001352D:D02 35550 3254 1/28/98 168 RTA00000177AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 135 RTA00000196AF.c.22.1 M00001352D:D02 35550 3256 1/28/98 359 RTA00000178R.l.08.1 M00001352D:D02 35550 3257 1/28/98 37 RTA00000178R.l.08.1 M00001383A:C03 39648 3259 1/28/98 337 RTA00000177AR.m.17.1 M00001358B:G10 14391 3259 1/28/98 361 RTA00000177AR.m.17.4 M00001355B:G10 14391 3259 1/28/98 361 RTA00000177AR.g.16.4 M00001355B:G10 14391 3260 1/28/98 361 RTA00000177AR.g.16.4 M00001347A:B10 13576 3261 1/28/98 36 RTA00000177AF.e.14.1 M00001354B:B0 11 13576 3262 1/28/98 48 RTA00000177AF.e.9.1 M00001347A:B10 13576 3263 1/28/98 48 RTA00000177AF.e.9.1 M00001343D:C04 37442 3264 2/24/98 308 RTA00000177AF.a.2.5 M0000133D:D60 4957 3265 1/28/98 164 RTA00000177AF.a.8.3 M00001350A:H01 7187 3266 1/28/98 283 RTA00000177AF.a.8.3 M00001350A:H01 7187 3267 1/28/98 383 RTA00000177AF.a.8.3 M00001350A:H01 7187 3268 1/28/98 383 RTA00000177AF.a.8.3 M00001350A:H01 7187 3270 1/28/98 296 RTA00000177AF.a.8.1 M00001350A:H01 7187 3271 1/28/98 297 RTA00000177AF.a.8.1 M00001350A:H01 7187 3272 1/28/98 37 RTA00000177AF.a.8.1 M00001350B:F06 4188 3277 1/28/98 37 RTA00000177AF.a.8.1 M00001350B:F06 4188 3277 1/28/98 37 RTA00000177AF.a.8.1 M00001350B:F06 4188 3277 1/28/98 37 RTA00000177AF.m.8.1 M00001350B:G10 14391 3276 1/28/98 37 RTA00000177AF.m.8.1 M00001350B:G10 14391 3276 1/28/98 37 RTA00000177AF.m.8.1 M00001350B:G10 14391 3277 1/28/98 37 RTA00000177AF.m.17.1 M00001355B:G10 14391 3278 1/28/98 37 RTA00000177AF.m.17.1 M00001355B:G10 14391 3279 1/28/98 37 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA		Appln	Appln			
3252 1/28/98 342 RTA00000196AF.d.10.1 M00001354C:B06 22256 3253 1/28/98 113 RTA00000177AR.k.23.1 M00001352D:D02 35550 3254 1/28/98 168 RTA00000177AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 270 RTA00000196AF.c.22.1 M00001352D:D02 35550 3256 1/28/98 359 RTA00000196AF.c.10.1 M00001352D:D02 35550 3258 1/28/98 359 RTA00000196AF.c.17.1 M00001348A:D04 12193 3259 1/28/98 337 RTA00000177AR.m.17.3 M00001358B:G10 14391 3259 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3260 1/28/98 361 RTA00000177AR.g.16.4 M00001347B:B10 13576 3261 1/28/98 265 RTA00000177AF.e.14.1 M00001343D:C04 37442 3263 1/28/98 308 RTA00000177AF.e.9.1 M00001343D:C04 37442 3263 1/28/98 164 RTA00000177AR.a.23.5 M00001349B:D07 32355 3264 1/28/98 164 RTA00000177AF.a.8.1 M00001349B:D07 32355 3266 1/28/98 15 RTA00000177AF.a.8.1 M00001349B:D07 32355 3267 1/28/98 283 RTA00000177AF.a.8.1 M00001347B:D00 49457 3266 1/28/98 15 RTA00000177AF.a.8.1 M00001347B:D00 49457 3267 1/28/98 296 RTA00000177AF.a.8.3 M00001356D:F06 4188 3268 1/28/98 383 RTA00000177AF.a.8.3 M00001356D:F06 4188 3269 1/28/98 296 RTA00000177AF.a.8.3 M00001356D:F06 4188 3269 1/28/98 39 RTA00000177AF.a.8.1 M00001355B:G10 14391 3271 1/28/98 297 RTA00000177AF.a.8.1 M00001355B:G10 14391 3272 1/28/98 30 RTA00000177AF.a.8.1 M00001356D:F06 4188 3273 1/28/98 30 RTA00000177AF.a.8.1 M00001355B:G10 14391 3276 1/28/98 30 RTA00000177AF.m.8.1 M00001355B:G10 14391 3277 1/28/98 30 RTA00000177AF.a.8.1 M00001355B:G10 14391 3278 1/28/98 30 RTA00000177AF.a.8.1 M00001355B:G10 14391 3279 1/28/98 30 RTA00000177AF.a.8.1 M00001355B:G10 14391 3279 1/28/98 30 RTA00000177AF.a.8.1 M00001355B:G10 14391 3279 1/28/98 33 RTA00000177AF.a.8.1 M00001355B:G10 14391 3280 1/28/98 33 RTA00000177AF.a.8.1 M00001355B:G10 14391 3280 1/28/98 33	3250	1/28/98	362			
3253 1/28/98 463 RTA00000177AR.k.23.4 M00001354B:B10 16934 3254 1/28/98 168 RTA00000177AR.k.23.4 M00001352D:D02 35550 3255 1/28/98 463 RTA00000177AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 168 RTA00000177AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 155 RTA0000017AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 270 RTA0000016AF.c.22.1 M00001352D:D02 35550 3255 1/28/98 379 RTA0000017AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 399 RTA0000017AR.k.23.1 M00001352D:D02 35550 3255 1/28/98 399 RTA0000017AR.h.17.1 M00001352D:D02 35550 3259 1/28/98 393 RTA0000017AR.m.17.3 M00001355B:G10 14391 3259 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3259 1/28/98 361 RTA00000177AR.m.17.4 M00001355B:G10 14391 3260 1/28/98 361 RTA00000177AF.e.14.1 M00001347A:B10 13576 3261 1/28/98 265 RTA00000177AF.e.9.1 M00001347A:B10 13576 3262 1/28/98 48 RTA00000177AF.e.9.1 M00001347A:B10 13576 3264 2/24/98 308 RTA00000177AR.a.23.5 M00001340:C04 37442 3263 1/28/98 164 RTA00000177AR.a.23.5 M00001339D:G02 6995 3264 2/24/98 308 RTA00000177AR.a.1.1 M00001350A:H01 7187 3266 1/28/98 15 RTA00000177AF.a.8.3 M00001350A:H01 7187 3267 1/28/98 383 RTA00000177AF.a.8.1 M00001350A:H01 7187 3267 1/28/98 383 RTA00000177AF.a.8.1 M00001350A:H01 7187 3268 1/28/98 383 RTA00000177AF.a.8.3 M00001350A:H01 7187 3271 1/28/98 296 RTA0000016AF.f.20.1 M00001350A:H01 7187 3271 1/28/98 297 RTA0000017AF.a.8.1 M00001350A:H01 7187 3272 1/28/98 330 RTA0000017AF.a.8.1 M00001350A:H01 7187 3273 1/28/98 39 RTA0000017AF.a.8.1 M00001350B:G10 4188 3276 1/28/98 33 RTA0000017AF.a.8.1 M00001350B:G10 4188 3277 1/28/98 39 RTA0000017AF.m.1.1 M00001355B:G10 14391 3276 1/28/98 33 RTA0000017AF.m.1.1 M00001355B:G10 14391 3277 1/28/98 33 RTA0000017AF.m.1.1 M00001355B:G10 14391 3278 1/28/98 33 RTA0000017AF.m.1.1 M00001355B:G10 14391 3279 1/28/98 33 RTA0000017AR.m.17.3 M00001355B:G10 14391 3279 1/28/98 33 RTA0000017AR.m.17.1 M00001355B:G10 14391 3279 1/28/98 33 RTA0000017AR.m.17.1 M00001355B:G10 14391 3280 1/28/98 33 RTA0000017AR.m.17.1 M00001355B:G10 14391 3281 1/28/98 33 RTA0000017AR.m.17.1 M00	3251	1/28/98	81			
3254 1/28/98 463 RTA00000177AR.k.23.4 M00001352D:D02 35550	3252	1/28/98	342	RTA00000196AF.d.10.1		
3254 1/28/98 168	3253	1/28/98	113			
3255 1/28/98 463 RTA00000177AR.k.23.4 M00001352D:D02 35550 32555 1/28/98 168 RTA00000177AR.k.23.1 M00001352D:D02 35550 32556 1/28/98 135 RTA00000196AF.c.22.1 M00001352D:D02 32550 32551 1/28/98 270 RTA00000178R.l.08.1 M00001352D:D02 32550 32558 1/28/98 359 RTA00000196AF.b.17.1 M00001355B:G10 41391 3259 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 41391 4	3254	1/28/98	463			
3255 1/28/98 168 RTA00000177AR.k.23.1 M00001352D:D02 35550 3256 1/28/98 135 RTA00000196AF.c.22.1 M00001352D:D02 22548 3257 1/28/98 359 RTA00000196AF.c.22.1 M00001383A:C03 39648 3258 1/28/98 359 RTA00000196AF.c.17.1 M00001383A:C03 39648 3258 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 14391 3259 1/28/98 330 RTA00000177AR.m.17.3 M00001355B:G10 14391 3260 1/28/98 361 RTA00000177AR.m.17.4 M00001355B:G10 14391 3260 1/28/98 265 RTA00000177AF.e.9.1 M00001347A:B10 13576 3261 1/28/98 56 RTA00000177AF.e.9.1 M00001347A:B10 3256 1/28/98 48 RTA00000177AF.e.9.1 M00001347D:D04 37442 3263 1/28/98 48 RTA00000177AF.a.23.5 M00001349D:G02 6995 3264 2/24/98 308 RTA00000177AF.a.23.5 M00001349D:G02 6995 3264 2/24/98 308 RTA00000177AF.a.8.3 M00001349D:G02 6995 3266 1/28/98 164 RTA00000193AR.i.14.4 M00004307C:A06 9457 3266 1/28/98 89 RTA00000177AF.a.8.3 M00001356D:F06 4188 3267 1/28/98 89 RTA00000177AF.a.8.3 M00001356D:F06 4188 3267 1/28/98 383 RTA00000177AF.a.8.3 M00001356D:F06 4188 3268 1/28/98 383 RTA00000177AF.a.8.3 M00001356D:F06 4188 3269 1/28/98 328 RTA00000177AF.a.8.3 M0000137D:G01 22774 3271 1/28/98 296 RTA00000178AF.f.20.3 M0000137D:G01 22774 3271 1/28/98 296 RTA00000178AF.f.20.3 M0000137D:G01 22774 3271 1/28/98 337 RTA00000177AF.m.8.1 M00001355B:G10 4391 3276 1/28/98 337 RTA00000177AF.m.8.1 M00001355B:G10 4391 3276 1/28/98 337 RTA00000177AF.m.8.1 M00001355B:G10 4391 3276 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3279 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3279 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3279 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3280 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3280 1/28/98 337 RTA00000	3254	1/28/98	168	RTA00000177AR.k.23.1		
3256 1/28/98 355 RTA00000196AF.c.22.1 M00001352D:C05 32548 3257 1/28/98 359 RTA000001778R.I.08.1 M00001383A:C03 39648 3258 1/28/98 359 RTA00000177AF.m.17.1 M00001348A:D04 12193 3259 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 14391 3259 1/28/98 330 RTA00000177AR.m.17.3 M00001355B:G10 14391 3259 1/28/98 361 RTA00000177AR.m.17.4 M00001355B:G10 14391 3260 1/28/98 361 RTA00000177AF.e.16.4 M00001347A:B10 13576 3261 1/28/98 365 RTA00000177AF.e.14.1 M00001343D:H07 23255 3262 1/28/98 48 RTA00000177AF.e.9.1 M00001343D:H07 23255 3263 1/28/98 48 RTA00000177AF.e.9.1 M00001343D:C04 37442 3263 1/28/98 48 RTA00000133AR.i.14.4 M00001343D:C04 37442 3265 1/28/98 164 RTA00000177AF.e.8.1 M00001350D:F06 4188 3267 1/28/98 283 RTA00000177AF.e.8.4 M00001350D:F06 4188 3267 1/28/98 38 RTA00000177AF.n.8.1 M00001350D:F06 4188 3268 1/28/98 38 RTA00000177AF.n.8.1 M00001350D:F06 4188 3268 1/28/98 38 RTA00000177AF.n.8.3 M00001350D:F06 4188 3269 1/28/98 38 RTA00000178AF.f.20.2 M00003847B:G03 0 3271 1/28/98 296 RTA00000178AF.f.20.3 M00001371C:E09 7172 3272 1/28/98 240 RTA00000178AF.e.1.6 M0000136D:F06 4188 3273 1/28/98 330 RTA00000177AF.m.8.1 M0000136D:F06 3274 1/28/98 330 RTA00000177AF.m.8.1 M0000135D:G10 3275 1/28/98 337 RTA00000177AF.m.8.1 M0000135D:G10 3276 1/28/98 337 RTA00000177AF.m.8.1 M0000135D:G10 34391 3276 1/28/98 337 RTA00000177AF.m.8.1 M0000135D:G10 43391 3276 1/28/98 337 RTA00000177AF.m.8.1 M0000135D:G10 43391 3276 1/28/98 337 RTA00000177AF.m.17.1 M0000135D:G10 43391 3279 1/28/98 337 RTA00000177AF.m.17.1 M0000135D:G10 43391 3279 1/28/98 337 RTA00000177AF.m.17.1 M0000135D:G10 4391 3279 1/28/98 337 RTA00000177AF.m.17.1 M0000135D:G10 4391 3280 1/28/98 337 RTA00000177AF.m.17.1 M0000135	3255	1/28/98	463	RTA00000177AR.k.23.4		
3257 1/28/98 270 RTA00000178R.I.08.1 M00001383A:C03 39648 3258 1/28/98 359 RTA00000196AF.b.17.1 M00001348A:D04 12193 3259 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 14391 3259 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3260 1/28/98 361 RTA00000177AR.m.17.4 M00001347A:B10 73255 3261 1/28/98 265 RTA00000177AF.e.9.1 M00001347A:B10 73442 3262 1/28/98 56 RTA00000177AF.e.9.1 M00001339D:G02 6995 3264 1/28/98 48 RTA000001535R.d.11.1 M00001339D:G02 6995 3264 1/28/98 164 RTA00000157AF.e.9.1 M00001330D:H08 0 3265 1/28/98 15 RTA00000177AF.n.8.2 M00001350A:H01 7187 3266 1/28/98 15 RTA00000177AF.n.8.1 M00001350A:H01 7187 3267 1/28/98 38	3255	1/28/98	168	RTA00000177AR.k.23.1		
3258 1/28/98 359 RTA00000196AF.b.17.1 M00001348A:D04 12193 3259 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3259 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3260 1/28/98 361 RTA00000177AR.m.17.4 M00001343D:H07 3261 3261 1/28/98 361 RTA00000177AR.e.14.1 M00001343D:H07 37442 3262 1/28/98 56 RTA00000177AF.e.9.1 M00001343D:C04 37442 3263 1/28/98 48 RTA00000177AF.e.9.1 M00001339D:G02 6995 3264 1/28/98 164 RTA00000177AF.e.9.1 M00001330D:G02 6995 3265 1/28/98 164 RTA0000017AF.e.8.4 M00001350A:H01 7187 3267 1/28/98 15 RTA0000017AF.e.8.4 M00001356D:F06 4188 3267 1/28/98 89 RTA0000017AF.e.8.3 M00000137D:G03 0 3268 1/28/98 132 <	3256	1/28/98	135	RTA00000196AF.c.22.1	M00001352D:C05	
1/28/98 493 RTA00000177AF.m.17.1 M00001355B:GI0 14391 3259 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:GI0 14391 3259 1/28/98 330 RTA00000177AR.m.17.3 M00001355B:GI0 14391 3259 1/28/98 361 RTA00000177AR.m.17.4 M00001355B:GI0 14391 3260 1/28/98 365 RTA00000177AR.e.16.4 M00001347A:BI0 13576 3261 1/28/98 56 RTA00000177AF.e.14.1 M00001343D:H07 32255 3262 1/28/98 48 RTA00000177AF.e.9.1 M00001334D:H07 32255 3264 2/24/98 308 RTA00000177AF.e.9.1 M00001339D:G02 6995 3264 2/24/98 308 RTA00000177AF.e.3.1 M00001330D:G02 6995 3266 1/28/98 283 RTA00000177AF.a.8.1 M00001350D:F06 4188 3267 1/28/98 89 RTA00000177AF.a.8.1 M00001350D:F06 4188 3267 1/28/98 89 RTA00000177AF.a.8.3 M00001350D:F06 4188 3268 1/28/98 333 RTA00000177AF.a.8.3 M00001350D:F06 4188 3269 1/28/98 332 RTA00000178AF.f.20.1 M00001371D:G01 22774 3271 1/28/98 296 RTA00000178AF.f.9.3 M00001371D:G01 22774 3271 1/28/98 240 RTA00000178AF.f.9.3 M00001371D:G01 22774 3271 1/28/98 340 RTA00000178AF.a.8.1 M00001360D:F06 32273 1/28/98 312 RTA00000178AF.a.8.1 M0000137D:G01 22774 3271 1/28/98 340 RTA00000178AF.a.8.1 M0000137D:G01 22774 3271 1/28/98 316 RTA00000178AF.a.8.1 M00001355B:G10 3275 1/28/98 330 RTA00000177AF.m.8.1 M00001355B:G10 4391 3276 1/28/98 337 RTA00000177AF.m.8.1 M00001355B:G10 4391 3276 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3278 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3279 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3279 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3279 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3280 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3280 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 4391 3280 1/28/98 337 RTA00000177AF.m.17.1	3257	1/28/98	270	RTA00000178R.1.08.1	M00001383A:C03	
1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 1439	3258	1/28/98	359	RTA00000196AF.b.17.1	M00001348A:D04	12193
3259 1/28/98 337	3259	1/28/98	493	RTA00000177AF.m.17.1	M00001355B:G10	14391
3259 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3260 1/28/98 361 RTA00000177AR.g.16.4 M00001347A:B10 13576 3261 1/28/98 56 RTA00000177AF.e.9.1 M00001343D:C04 37442 3263 1/28/98 48 RTA00000177AR.a.23.5 M00001339D:G02 6995 3264 2/24/98 308 RTA00000173AR.a.23.5 M00001339D:G02 6995 3266 1/28/98 164 RTA00000193AR.i.14.4 M00004307C:A06 9457 3266 1/28/98 15 RTA00000177AR.n.8.1 M00001350A:H01 7187 3267 1/28/98 15 RTA00000177AR.n.8.1 M00001350D:F06 4188 3268 1/28/98 383 RTA00000178AF.g.20.2 M00003847B:G03 0 3269 1/28/98 132 RTA00000178AF.g.20.1 M0000137C:CF07 39881 3270 1/28/98 296 RTA00000178AF.g.20.1 M0000137C:CE09 7172 3272 1/28/98 296			337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3260 1/28/98 361 RTA00000177AR.g.16.4 M00001347A:B10 13576 3261 1/28/98 265 RTA00000177AF.e.14.1 M00001343D:H07 23255 3262 1/28/98 56 RTA00000177AF.e.19.1 M00001343D:H02 37442 3263 1/28/98 48 RTA00000177AF.a.23.5 M00001339D:G02 6995 3264 2/24/98 308 RTA00000133R.i.14.4 M00004692A:H08 0 3265 1/28/98 164 RTA0000017AF.i.8.4 M00001350A:H01 7187 3266 1/28/98 15 RTA0000017AF.i.8.4 M00001350D:F06 4188 3267 1/28/98 15 RTA0000017AF.n.8.3 M00001356D:F06 4188 3268 1/28/98 38 RTA0000017AF.f.20.2 M000003847B:G03 0 3270 1/28/98 132 RTA0000017AF.f.20.1 M0000137IC:E07 39881 3271 1/28/98 296 RTA0000017AF.f.20.1 M0000137IC:E07 39881 3272 1/28/98 12 RT		1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3261 1/28/98 265 RTA00000177AF.e.14.1 M00001343D:H07 23255 3262 1/28/98 56 RTA00000177AF.e.9.1 M00001343D:C04 37442 3263 1/28/98 48 RTA00000177AR.a.23.5 M0000139D:G02 6995 3264 2/24/98 308 RTA00000153Rd.i.1.1 M00004692A:H08 0 3265 1/28/98 164 RTA00000177AF.i.8.4 M00001350A:H01 7187 3266 1/28/98 283 RTA00000177AF.i.8.4 M00001356D:F06 4188 3267 1/28/98 15 RTA00000177AF.i.8.3 M00001356D:F06 4188 3268 1/28/98 383 RTA00000178AF.f.20.2 M00003847B:G03 0 3269 1/28/98 132 RTA00000178AF.f.20.1 M00001371C:E07 39881 3270 1/28/98 296 RTA00000178AF.f.20.1 M00001371C:E09 7172 3272 1/28/98 240 RTA00000177AF.m.8.1 M00001360D:E1 2074 3273 1/28/98 16 R				RTA00000177AR.g.16.4	M00001347A:B10	13576
3262 1/28/98 56 RTA00000177AF.e.9.1 M00001343D:C04 37442 3263 1/28/98 48 RTA00000177AR.a.23.5 M0000133DD:G02 6995 3264 2/24/98 308 RTA00000173AR.a.23.5 M00004692A:H08 0 3265 1/28/98 164 RTA00000177AF.is.4 M00004307C:A06 9457 3266 1/28/98 283 RTA00000177AF.is.4 M00001350A:H01 7187 3267 1/28/98 15 RTA00000177AF.n.8.1 M00001356D:F06 4188 3267 1/28/98 89 RTA00000177AF.n.8.3 M00001356D:F06 4188 3268 1/28/98 383 RTA00000196AF.f.20.2 M00003847B:G03 0 3270 1/28/98 132 RTA00000178AF.f.20.1 M00001371D:G01 22774 3271 1/28/98 296 RTA00000178AF.f.20.1 M00001369A:H12 2664 3273 1/28/98 297 RTA0000017AF.e.1.1 M00001369A:H12 2664 3273 1/28/98 16 RTA0					M00001343D:H07	23255
3263 1/28/98 48 RTA00000177AR.a.23.5 M00001339D:G02 6995 3264 2/24/98 308 RTA00000353R.d.11.1 M00004692A:H08 0 3265 1/28/98 164 RTA0000017AR.i.14.4 M00004307C:A06 9457 3266 1/28/98 283 RTA0000017AR.ii.8.4 M00001350A:H01 7187 3267 1/28/98 89 RTA0000017AR.ii.8.1 M00001356D:F06 4188 3268 1/28/98 89 RTA0000017AF.ii.8.3 M00001376D:F06 4188 3269 1/28/98 89 RTA0000017AF.ii.8.3 M00001370C:F07 39881 3269 1/28/98 132 RTA0000017AF.ii.8.3 M00001371D:G01 22774 3271 1/28/98 296 RTA0000017AF.f.20.3 M00001371D:G01 22774 3271 1/28/98 297 RTA0000017AF.ei.1.1 M00001369A:H12 2664 3272 1/28/98 16 RTA0000017AF.ei.1.1 M00001356D:H02 8010 3274 1/28/98 15 RT					M00001343D:C04	37442
3264 2/24/98 308 RTA00000353R.d.11.1 M00004692A:H08 0 3265 1/28/98 164 RTA00000193AR.i.14.4 M00004307C:A06 9457 3266 1/28/98 283 RTA00000177AR.i.8.4 M00001350D:F06 4188 3267 1/28/98 89 RTA00000177AR.i.8.1 M00001356D:F06 4188 3268 1/28/98 383 RTA00000177AF.ii.8.3 M00001356D:F06 4188 3269 1/28/98 132 RTA00000178AF.f20.2 M00003847B:G03 0 3270 1/28/98 296 RTA00000178AF.f20.1 M00001371D:G01 22774 3271 1/28/98 297 RTA00000178AF.f.20.1 M00001371C:E09 7172 3272 1/28/98 16 RTA00000178AF.e.1.1 M00001369A:H12 2664 3273 1/28/98 16 RTA00000178AF.e.1.1 M00001354C:C10 8010 3275 1/28/98 15 RTA00000177AF.m.8.1 M00001355B:G10 4191 3276 1/28/98 330 RTA				RTA00000177AR.a.23.5	M00001339D:G02	6995
3265 1/28/98 164 RTA00000193AR.i.14.4 M00004307C:A06 9457 3266 1/28/98 283 RTA00000177AF.i.8.4 M00001350A:H01 7187 3267 1/28/98 89 RTA00000177AF.i.8.3 M00001356D:F06 4188 3267 1/28/98 89 RTA00000177AF.i.8.3 M00001356D:F06 4188 3268 1/28/98 383 RTA00000177AF.i.8.3 M00001356D:F06 4188 3269 1/28/98 312 RTA00000178AF.f.20.3 M0000137D:G01 22774 3270 1/28/98 296 RTA00000178AF.f.20.1 M00001371D:G01 22774 3271 1/28/98 240 RTA00000178AF.e.1.1 M00001369A:H12 2664 3272 1/28/98 16 RTA00000177AF.m.8.1 M00001354C:C10 8010 3275 1/28/98 154 RTA00000177AF.m.8.1 M00001360D:E11 1039 3276 1/28/98 33 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 37					M00004692A:H08	0
3266 1/28/98 283 RTA00000177AF.i.8.4 M00001350A:H01 7187 3267 1/28/98 15 RTA00000177AR.n.8.1 M00001356D:F06 4188 3267 1/28/98 89 RTA00000177AF.n.8.3 M00001356D:F06 4188 3268 1/28/98 383 RTA0000019F.f.20.2 M00003847B:G03 0 3269 1/28/98 132 RTA00000178AF.f.20.1 M00001371D:G01 22774 3271 1/28/98 296 RTA00000178AF.f.20.1 M00001371D:G01 22774 3271 1/28/98 297 RTA00000178AF.f.9.3 M00001371C:E09 7172 3272 1/28/98 240 RTA00000178AF.e.1.1 M00001369A:H12 2664 3273 1/28/98 16 RTA0000017AF.m.8.1 M00001354C:H02 39252 3274 1/28/98 154 RTA0000017AF.m.8.1 M00001360D:E11 1039 3276 1/28/98 330 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 37					M00004307C:A06	9457
3267 1/28/98 15 RTA00000177AR.n.8.1 M00001356D:F06 4188 3267 1/28/98 89 RTA00000177AR.n.8.3 M00001356D:F06 4188 3268 1/28/98 383 RTA00000178AF.c.0.2 M00003847B:G03 0 3269 1/28/98 132 RTA00000178AF.c.0.3 M0000137C:F07 39881 3270 1/28/98 296 RTA00000178AF.c.0.1 M00001371D:G01 22774 3271 1/28/98 297 RTA00000178AF.c.1.1 M00001371D:E09 7172 3272 1/28/98 240 RTA00000178AF.c.1.1 M00001369A:H12 2664 3273 1/28/98 16 RTA00000017AF.m.8.1 M00001354C:C10 8010 3274 1/28/98 112 RTA00000177AF.m.8.1 M00001354C:C10 8010 3275 1/28/98 330 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 14391 3277 1/28/98 35 <td< td=""><td></td><td></td><td></td><td></td><td>M00001350A:H01</td><td>7187</td></td<>					M00001350A:H01	7187
3267 1/28/98 89 RTA00000177AF.n.8.3 M00001356D:F06 4188 3268 1/28/98 383 RTA00000199F.f.20.2 M00003847B:G03 0 3269 1/28/98 132 RTA00000178AF.f.20.3 M00001371D:G01 22774 3270 1/28/98 296 RTA00000178AF.f.20.1 M00001371D:G01 22774 3271 1/28/98 297 RTA00000178AF.e.13. M00001371C:E09 7172 3272 1/28/98 240 RTA00000178AF.e.16.1 M00001369A:H12 2664 3273 1/28/98 16 RTA00000177AF.m.8.1 M00001363C:H02 39252 3274 1/28/98 112 RTA00000177AF.m.8.1 M00001355B:G10 8010 3275 1/28/98 154 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 330 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AF.m.8.3 M00001355B:G10 14391 3277 1/28/98 337					M00001356D:F06	4188
3268 1/28/98 383 RTA00000199F.f.20.2 M00003847B:G03 0 3269 1/28/98 132 RTA00000178AF.f.20.3 M00001372C:F07 39881 3270 1/28/98 296 RTA00000178AF.f.20.1 M00001371D:G01 22774 3271 1/28/98 297 RTA00000178AF.f.9.3 M00001369A:H12 2664 3272 1/28/98 16 RTA00000178AF.e.1.1 M00001369A:H12 2664 3273 1/28/98 16 RTA00000177AF.m.8.1 M00001363C:H02 39252 3274 1/28/98 112 RTA00000177AF.m.8.1 M00001364C:C10 8010 3275 1/28/98 154 RTA00000177AF.m.8.1 M00001360D:E11 1039 3276 1/28/98 330 RTA00000177AF.m.17.4 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AF.m.8.3 M00001355B:G10 14391 3277 1/28/98 89 RTA00000177AF.n.8.3 M00001356D:F06 4188 3278 1/28/98 493					M00001356D:F06	4188
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3270 1/28/98 296 RTA00000196AF.f.20.1 M00001371D:G01 22774 3271 1/28/98 297 RTA00000178AF.f.9.3 M00001371C:E09 7172 3272 1/28/98 240 RTA00000178AF.e.1.1 M00001369A:H12 2664 3273 1/28/98 16 RTA00000196AF.e.16.1 M00001363C:H02 39252 3274 1/28/98 112 RTA00000177AF.m.8.1 M00001354C:C10 8010 3275 1/28/98 154 RTA00000177AF.m.8.1 M00001355B:G10 14391 3276 1/28/98 330 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AF.n.8.3 M00001355B:G10 14391 3276 1/28/98 493 RTA00000177AF.n.8.1 M00001355B:G10 14391 3276 1/28/98 493 RTA00000177AF.n.8.1 M00001355B:G10 14381 3279 1/28/98 337 <td></td> <td></td> <td></td> <td></td> <td>M00001372C:F07</td> <td>39881</td>					M00001372C:F07	39881
3271 1/28/98 297 RTA00000178AF.f.9.3 M00001371C:E09 7172 3272 1/28/98 240 RTA00000178AF.e.1.1 M00001369A:H12 2664 3273 1/28/98 16 RTA00000196AF.e.16.1 M00001354C:C10 8010 3274 1/28/98 112 RTA00000177AF.m.8.1 M00001354C:C10 8010 3275 1/28/98 330 RTA00000177AF.m.8.1 M00001360D:E11 1039 3276 1/28/98 330 RTA00000177AF.m.17.4 M00001355B:G10 14391 3276 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AF.m.17.3 M00001355B:G10 14391 3277 1/28/98 89 RTA00000177AF.m.8.3 M00001356D:F06 4188 3278 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AF.m.17.1 M00001355B:G10 14391 3279 1/28/98 337						22774
3272 1/28/98 240 RTA00000178AF.e.1.1 M00001369A:H12 2664 3273 1/28/98 16 RTA00000196AF.e.16.1 M00001363C:H02 39252 3274 1/28/98 112 RTA00000177AF.m.8.1 M00001354C:C10 8010 3275 1/28/98 154 RTA00000177AF.m.8.1 M00001360D:E11 1039 3276 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3276 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AF.m.17.3 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AF.m.8.3 M00001355B:G10 14391 3277 1/28/98 89 RTA00000177AF.m.17.1 M00001355B:G10 4188 3278 1/28/98 15 RTA00000177AF.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 <td></td> <td></td> <td></td> <td></td> <td>M00001371C:E09</td> <td>7172</td>					M00001371C:E09	7172
3273 1/28/98 16 RTA00000196AF.e.16.1 M00001363C:H02 39252 3274 1/28/98 112 RTA00000177AF.m.8.1 M00001354C:C10 8010 3275 1/28/98 154 RTA00000196F.e.7.1 M00001360D:E11 1039 3276 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3276 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3277 1/28/98 89 RTA00000177AR.m.8.3 M00001356D:F06 4188 3278 1/28/98 493 RTA00000177AR.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3279 1/28/98 337 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 </td <td></td> <td></td> <td></td> <td></td> <td>M00001369A:H12</td> <td>2664</td>					M00001369A:H12	2664
3274 1/28/98 112 RTA00000177AF.m.8.1 M00001354C:C10 8010 3275 1/28/98 154 RTA00000196F.e.7.1 M00001360D:E11 1039 3276 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3276 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3277 1/28/98 89 RTA00000177AR.m.8.3 M00001356D:F06 4188 3277 1/28/98 15 RTA00000177AR.m.8.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3279 1/28/98 337 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 <td></td> <td></td> <td></td> <td></td> <td>M00001363C:H02</td> <td>39252</td>					M00001363C:H02	39252
3275 1/28/98 154 RTA00000196F.e.7.1 M00001360D:E11 1039 3276 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3276 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3277 1/28/98 89 RTA00000177AF.n.8.3 M00001356D:F06 4188 3277 1/28/98 15 RTA00000177AR.n.8.1 M00001356D:F06 4188 3278 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3278 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 337 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3281 1/28/98 337 </td <td></td> <td></td> <td></td> <td>RTA00000177AF.m.8.1</td> <td>M00001354C:C10</td> <td>8010</td>				RTA00000177AF.m.8.1	M00001354C:C10	8010
3276 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3276 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3277 1/28/98 89 RTA00000177AF.n.8.3 M00001356D:F06 4188 3277 1/28/98 15 RTA00000177AR.n.8.1 M00001356D:F06 4188 3278 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3278 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3281 1/28/98 33				RTA00000196F.e.7.1	M00001360D:E11	1039
3276 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3276 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3277 1/28/98 89 RTA00000177AF.n.8.3 M00001356D:F06 4188 3277 1/28/98 15 RTA00000177AR.n.8.1 M00001356D:F06 4188 3278 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3278 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 33			330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3276 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3277 1/28/98 89 RTA00000177AF.n.8.3 M00001356D:F06 4188 3277 1/28/98 15 RTA00000177AR.n.8.1 M00001356D:F06 4188 3278 1/28/98 493 RTA00000177AR.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3278 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3280 1/28/98 337 RTA00000177AR.m.17.1 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 33				RTA00000177AF.m.17.1	M00001355B:G10	14391
3277 1/28/98 89 RTA00000177AF.n.8.3 M00001356D:F06 4188 3277 1/28/98 15 RTA00000177AF.n.8.1 M00001356D:F06 4188 3278 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3278 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AF.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 33		1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3277 1/28/98 15 RTA00000177AR.n.8.1 M00001356D:F06 4188 3278 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3278 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 <td< td=""><td></td><td></td><td>89</td><td>RTA00000177AF.n.8.3</td><td>M00001356D:F06</td><td>4188</td></td<>			89	RTA00000177AF.n.8.3	M00001356D:F06	4188
3278 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3278 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.3 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 337 RTA00000177AR.m.17.1 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3282 1/28/98		1/28/98	15	RTA00000177AR.n.8.1	M00001356D:F06	4188
3278 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3278 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3280 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98		1/28/98	493	RTA00000177AF.m.17.1	M00001355B:G10	14391
3279 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000177AR.m.17.4 M00001355B:G10 14391 4490 RTA000000177AR.m.17.4		1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3279 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3279 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685	3278	1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3279 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 330 RTA00000177AF.m.17.4 M00001355B:G10 14391 3280 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685	3279	1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3280 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3280 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685		1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3280 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3280 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685	3279	1/28/98	493	RTA00000177AF.m.17.1	M00001355B:G10	14391
3280 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685	3280	1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685	3280	1/28/98	493	RTA00000177AF.m.17.1		
3281 1/28/98 337 RTA00000177AR.m.17.3 M00001355B:G10 14391 3281 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685		1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3281 1/28/98 493 RTA00000177AF.m.17.1 M00001355B:G10 14391 3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685				RTA00000177AR.m.17.3	M00001355B:G10	14391
3281 1/28/98 330 RTA00000177AR.m.17.4 M00001355B:G10 14391 3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685		1/28/98	493	RTA00000177AF.m.17.1		
3282 1/28/98 169 RTA00000196AF.g.24.1 M00001380C:F02 8685			330	RTA00000177AR.m.17.4	M00001355B:G10	
		1/28/98	169			
	3283	1/28/98	363	RTA00000196AF.e.14.1	M00001362C:A10	12850

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
3284	1/28/98	92	RTA00000198AF.j.18.1	M00001653B:G07	22759
3284	1/28/98	433	RTA00000198R.j.18.1	M00001653B:G07	22759
3285	1/28/98	537	RTA00000188AF.g.14.1	M00003774C:D02	0
3286	1/28/98	434	RTA00000187AR.d.2.2	M00001664C:H10	4892
3287	1/28/98	70 3	RTA00000198F.1.09.1	M00001664B:D06	3611
3288	1/28/98	430	RTA00000198R.k.23.1	M00001661B:C08	8995
3288	1/28/98	294	RTA00000198AF.k.23.1	M00001661B:C08	8995
3289	1/28/98	294	RTA00000198AF.k.23.1	M00001661B:C08	8995
3289	1/28/98	430	RTA00000198R.k.23.1	M00001661B:C08	8995
3290	1/28/98	754	RTA00000187AF.I.11.1	M00001681A:F03	4482
3291	1/28/98	732	RTA00000186AF.p.01.2	M00001654D:G11	40440
3292	1/28/98	475	RTA00000187AR.m.3.3	M00001682C:B12	17055
3293	1/28/98	433	RTA00000198R.j.18.1	M00001653B:G07	22759
3293	1/28/98	92	RTA00000198AF.j.18.1	M00001653B:G07	22759
3294	1/28/98	555	RTA00000198AF.j.08.1	M00001651B:A11	10983
3295	1/28/98	399	RTA00000186AF.m.15.2	M00001649C:B10	40122
3296	1/28/98	575	RTA00000186AF.I.12.2	M00001645A:C12	19267
3297	1/28/98	666	RTA00000198F.i.10.1	M00001640B:F03	12792
3298	1/28/98	654	RTA00000186AF.j.21.2	M00001639D:B07	22506
3299	1/28/98	670	RTA00000186AF.p.17.3	M00001656B:A07	38383
3300	1/28/98	393	RTA00000188AF.b.14.1	M00003754D:D02	0
3301	1/28/98	422	RTA00000189AF.b.12.1	M00003829B:G03	17233
3301	1/28/98	210	RTA00000189AR.b.12.1	M00003829B:G03	17233
3302	1/28/98	587	RTA00000199F.a.3.1	M00003772D:E10	16617
3303	1/28/98	394	RTA00000198AF.p.22.1	M00003771A:G10	0
3304	1/28/98	542	RTA00000198R.p.12.1	M00003763D:E10	8878
3304	1/28/98	64	RTA00000198AF.p.12.1	M00003763D:E10	8878
3305	1/28/98	64	RTA00000198AF.p.12.1	M00003763D:E10	8878
3305	1/28/98	542	RTA00000198R.p.12.1	M00003763D:E10	8878
3306	1/28/98	465	RTA00000187AF.k.20.1	M00001680B:C01	3648
3307	1/28/98	423	RTA00000188AR.b.17.1	M00003755A:B03	10662
3308	1/28/98	711	RTA00000198F.i.2.1	M00001637B:E07	8076
3309	1/28/98	497	RTA00000198AF.o.09.1	M00003751B:A05	4310
3309	1/28/98	506	RTA00000198R.o.09.1	M00003751B:A05	4310
3310	1/28/98	506	RTA00000198R.o.09.1	M00003751B:A05	4310
3310	1/28/98	497	RTA00000198AF.o.09.1	M00003751B:A05	4310
3311	1/28/98	432	RTA00000198AF.o.05.1	M00003750A:D01	26702
3311	1/28/98	49	RTA00000198R.o.05.1	M00003750A:D01	26702
3312	1/28/98	49	RTA00000198R.o.05.1	M00003750A:D01	26702
3312	1/28/98	432	RTA00000198AF.o.05.1	M00003750A:D01	26702
3313	1/28/98	585	RTA00000198AF.n.18.1	M00001771A:A07	16715
3314	1/28/98	527	RTA00000198R.m.23.1	M00001684B:G03	38469
3315	1/28/98	471	RTA00000188AF.e.2.1	M00003763B:H01	0
3316	1/28/98	171	RTA00000198R.c.07.1	M00001575D:G05	19181
3316	1/28/98	525	RTA00000198AF.c.7.1	M00001575D:G05	19181
3317	1/28/98	557	RTA00000198AF.d.9.1	M00001587D:A10	8841
3318	1/28/98	523	RTA00000198AF.d.4.1	M00001586D:E02	22435
3319	1/28/98	441	RTA00000185AF.e.6.1	M00001583B:E10	0
3320	1/28/98	439	RTA00000185AF.d.14.2	M00001579D:G07	8071
3321	1/28/98	561	RTA00000185AR.d.10.1	M00001579C:H10	0

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
110.	Priority	Priority			11.5
	Appln	Appln			
3322	1/28/98	277	RTA00000198AR.i.08.1	M00001639A:F10	9807
3322	1/28/98	487	RTA00000198F.i.8.1	M00001639A:F10	9807
3323	1/28/98	525	RTA00000198AF.c.7.1	M00001575D:G05	19181
3323	1/28/98	171	RTA00000198R.c.07.1	M00001575D:G05	19181
3324	2/24/98	317	RTA00000195AF.b.21.1	M00001595B:A09	39055
3324	1/28/98	602	RTA00000195AF.b.21.1	M00001595B:A09	39055
3325	1/28/98	507	RTA00000198AF.c.5.1	M00001573D:F10	53802
3326	1/28/98	414	RTA00000185AR.b.15.1	M00001573D:F04	39813
3326	1/28/98	428	RTA00000185AF.b.15.2	M00001573D:F04	39813
3327	1/28/98	428	RTA00000185AF.b.15.2	M00001573D:F04	39813
3327	1/28/98	414	RTA00000185AR.b.15.1	M00001573D:F04	39813
3328	1/28/98	414	RTA00000185AR.b.15.1	M00001573D:F04	39813
3328	1/28/98	428	RTA00000185AF.b.15.2	M00001573D:F04	39813
3329	1/28/98	428	RTA00000185AF.b.15.2	M00001573D:F04	39813
3329	1/28/98	414	RTA00000185AR.b.15.1	M00001573D:F04	39813
3330	1/28/98	392	RTA00000185AF.b.11.2	M00001573C:D03	9024
3331	1/28/98	549	RTA00000198AF.c.16.1	M00001579C:B11	26801
3332	1/28/98	628	RTA00000198AF.g.16.1	M00001621D:D03	6602
3333	1/28/98	616	RTA00000188AF.m.07.1	M00003798D:E03	23183
3334	1/28/98	489	RTA00000186AF.h.22.1	M00001634B:C10	16485
3335	1/28/98	655	RTA00000186AF.g.8.2	M00001630B:A11	8065
3336	1/28/98	592	RTA00000186AF.e.18.1	M00001624C:A06	0
3337	1/28/98	713	RTA00000198AF.g.21.1	M00001624A:F09	6273
3338	1/28/98	554	RTA00000186AR.e.07.4	M00001623D:G03	4175
3338	1/28/98	400	RTA00000186AR.e.07.3	M00001623D:G03	4175
3339	1/28/98	467	RTA00000195AF.b.19.1	M00001589A:D12	77678
3340	1/28/98	646	RTA00000186AF.d.24.1	M00001623C:H07	3114
3341	1/28/98	740	RTA00000198AF.d.15.1	M00001590C:H08	5997
3342	1/28/98	504	RTA00000198AF.g.2.1	M00001615C:D02	16640
3343	1/28/98	470	RTA00000198AF.f.16.1	M00001614A:E06	0
3344	1/28/98	388	RTA00000185AF.n.17.1	M00001609B:A11	5336
3345	1/28/98	495	RTA00000185AF.j.21.1	M00001597A:E12	0
3346	2/24/98	317	RTA00000195AF.b.21.1	M00001595B:A09	39055
3346	1/28/98	602	RTA00000195AF.b.21.1	M00001595B:A09	39055
3347	1/28/98	487	RTA00000198F.i.8.1	M00001639A:F10	9807
3347	1/28/98	277	RTA00000198AR.i.08.1	M00001639A:F10	9807
3348	1/28/98	554	RTA00000186AR.e.07.4	M00001623D:G03	4175
3348	1/28/98	400	RTA00000186AR.e.07.3	M00001623D:G03	4175
3349	1/28/98	699	RTA00000178AF.a.12.1	M00001362B:H06	0
3350	1/28/98	416	RTA00000199F.a.5.1	M00003773B:G01	22134
3351	1/28/98	656	RTA00000178AR.h.22.3	M00001376B:A08	19230
3351	1/28/98	657	RTA00000178AR.h.22.2	M00001376B:A08	19230
3351	2/24/98	1137	RTA00000345F.d.03.1	M00001376B:A08	19230
3352	1/28/98	656	RTA00000178AR.h.22.3	M00001376B:A08	19230
3352	1/28/98	6 57	RTA00000178AR.h.22.2	M00001376B:A08	19230
3352	2/24/98	1137	RTA00000345F.d.03.1	M00001376B:A08	19230
3353	1/28/98	522	RTA00000178AR.h.17.2	M00001376A:C05	23824
3353	2/24/98	1095	RTA00000345F.c.12.1	M00001376A:C05	23824
3354	1/28/98	522	RTA00000178AR.h.17.2	M00001376A:C05	23824
3354	2/24/98	1095	RTA00000345F.c.12.1	M00001376A:C05	23824

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
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3355	1/28/98	656	RTA00000178AR.h.22.3	M00001376B:A08	19230
3355	1/28/98	657	RTA00000178AR.h.22.2	M00001376B:A08	19230
3355	2/24/98	1137	RTA00000345F.d.03.1	M00001376B:A08	19230
3356	1/28/98	566	RTA00000195F.a.4.1	M00001372C:G12	20470
3357	1/28/98	657	RTA00000178AR.h.22.2	M00001376B:A08	19230
3357	1/28/98	656	RTA00000178AR.h.22.3	M00001376B:A08	19230
3357	2/24/98	1137	RTA00000345F.d.03.1	M00001376B:A08	19230
3358	1/28/98	605	RTA00000196F.e.9.1	M00001361A:H07	23300
3359	1/28/98	532	RTA00000177AF.o.4.1	M00001358C:C06	0
3360	1/28/98	493	RTA00000177AF.m.17.1	M00001355B:G10	14391
3360	1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3360	1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3361	1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3361	1/28/98	493	RTA00000177AF.m.17.1	M00001355B:G10	14391
3361	1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3362	1/28/98	337	RTA00000177AR.m.17.3	M00001355B:G10	14391
3362	1/28/98	493	RTA00000177AF.m.17.1	M00001355B:G10	14391
3362	1/28/98	330	RTA00000177AR.m.17.4	M00001355B:G10	14391
3363	1/28/98	742	RTA00000177AF.m.1.1	M00001353D:D10	14929
3364	1/28/98	547	RTA00000196AF.g.8.1	M00001375B:G12	39665
3365	1/28/98	510	RTA00000178AF.n.23.1	M00001387B:E02	3298
3366	1/28/98	606	RTA00000179AR.e.01.4	M00001395A:C09	2493
3367	2/24/98	1065	RTA00000195R.a.06.1	M00001394A:E04	35265
3367	1/28/98	595	RTA00000195R.a.06.1	M00001394A:E04	35265
3368	2/24/98	1065	RTA00000195R.a.06.1	M00001394A:E04	35265
3368	1/28/98	595	RTA00000195R.a.06.1	M00001394A:E04	35265
3369	1/28/98	370	RTA00000179AF.c.15.3	M00001392D:H06	2995
3369	1/28/98	460	RTA00000179AF.c.15.1	M00001392D:H06	2995
3370	1/28/98	370	RTA00000179AF.c.15.3	M00001392D:H06	2995
3370	1/28/98	460	RTA00000179AF.c.15.1	M00001392D:H06	2995
3371	1/28/98	657	RTA00000178AR.h.22.2	M00001376B:A08	19230
3371	1/28/98	656	RTA00000178AR.h.22.3	M00001376B:A08	19230
3371	2/24/98	1137	RTA00000345F.d.03.1	M00001376B:A08	19230
3372	1/28/98	675	RTA00000179AR.b.21.3	M00001392C:D05	4366
3372	2/24/98	1264	RTA00000345F.e.13.1	M00001392C:D05	4366
3373	1/28/98	168	RTA00000177AR.k.23.1	M00001352D:D02	35550
3373	1/28/98	463	RTA00000177AR.k.23.4	M00001352D:D02	35550
3374	1/28/98	652	RTA00000178AR.m.21.4	M00001385A:F12	7861
3374	1/28/98	653	RTA00000178AR.m.21.5	M00001385A:F12	7861
3375	1/28/98	653	RTA00000178AR.m.21.5	M00001385A:F12	7861
3375	1/28/98	652	RTA00000178AR.m.21.4	M00001385A:F12	7861
3376	1/28/98	672	RTA00000196AF.h.09.1	M00001382B:F12	8015
3377	1/28/98	668	RTA00000178AF.i.17.1	M00001377C:E12	0
3378	1/28/98	746	RTA00000178AF,i.01.2	M00001376B:F03	4
3379	1/28/98	656	RTA00000178AR.h.22.3	M00001376B:A08	19230
3379	1/28/98	657	RTA00000178AR.h.22.2	M00001376B:A08	19230
3379	2/24/98	1137	RTA00000345F.d.03.1	M00001376B:A08	19230
3380	1/28/98	675	RTA00000179AR.b.21.3	M00001392C:D05	4366
3380	2/24/98	1264	RTA00000345F.e.13.1	M00001392C:D05	4366
3381	1/28/98	651	RTA00000189AR.d.22.2	M00003844C:B11	6539

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
3382	1/28/98	444	RTA00000189AF.I.16.1	M00003879A:G05	0
3383	1/28/98	648	RTA00000199F.i.9.1	M00003878C:E04	7
3384	2/24/98	678	RTA00000195AF.c.24.1	M00003860D:H07	0
3384	1/28/98	412	RTA00000195AF.c.24.1	M00003860D:H07	0
3385	1/28/98	412	RTA00000195AF.c.24.1	M00003860D:H07	0
3385	2/24/98	678	RTA00000195AF.c.24.1	M00003860D:H07	0
3386	1/28/98	484	RTA00000199F.g.20.2	M00003860D:A01	15767
3387	1/28/98	398	RTA00000177AR.I.13.3	M00001353A:G12	8078
3388	1/28/98	556	RTA00000199F.f.17.2	M00003845D:B04	22905
3389	1/28/98	545	RTA00000196F.a.2.1	M00001338B:E02	3575
3390	1/28/98	406	RTA00000199F.f.09.2	M00003842B:D09	22907
3390	1/28/98	78	RTA00000199R.f.09.1	M00003842B:D09	22907
3391	1/28/98	78	RTA00000199R.f.09.1	M00003842B:D09	22907
3391	1/28/98	406	RTA00000199F.f.09.2	M00003842B:D09	22907
3392	1/28/98	692	RTA00000199F.e.4.1	M00003820B:C05	0
3393	1/28/98	458	RTA00000199R.d.16.1	M00003812C:A05	24191
3394	1/28/98	755	RTA00000199F.c.21.2	M00003803C:D09	5070
3395	1/28/98	505	RTA00000188AF.n.03.1	M00003801B:B10	9443
3396	1/28/98	714	RTA00000199R.g.07.1	M00003853D:D03	0
3397	1/28/98	724	RTA00000177AR.f.15.4	M00001345B:E10	9062
3398	1/28/98	623	RTA00000198R.b.24.1	M00001571D:B11	19047
3398	1/28/98	748	RTA00000198AF.b.24.1	M00001571D:B11	19047
3399	1/28/98	395	RTA00000196R.c.21.2	M00001352C:H10	0
3400	1/28/98	593	RTA00000196R.c.14.2	M00001352B:F04	23105
3400	1/28/98	48 5	RTA00000196AF.c.14.1	M00001352B:F04	23105
3401	1/28/98	485	RTA00000196AF.c.14.1	M00001352B:F04	23105
3401	1/28/98	593	RTA00000196R.c.14.2	M00001352B:F04	23105
3402	1/28/98	645	RTA00000177AF.k.9.1	M00001352A:E02	16245
3403	1/28/98	576	RTA00000196AF.c.7.1	M00001350B:G11	0
3404	1/28/98	737	RTA00000189AR.m.9.1	M00003880B:C08	2917
3405	1/28/98	728	RTA00000177AR.f.17.4	M00001345C:B01	8594
3406	1/28/98	453	RTA00000199AF.i.20.1	M00003881A:D09	9544
3407	1/28/98	440	RTA00000177AR.f.13.4	M00001345A:G11	10480
3408	1/28/98	680	RTA00000177AF.f.10.1	M00001345A:E01	6420
3409	1/28/98	573	RTA00000196AF.b.7.1	M00001344A:G07	7774
3410	1/28/98	402	RTA00000177AF.b.21.4	M00001341A:F12	4443
3411	1/28/98	702	RTA00000177AR.b.8.5	M00001340B:A06	17062
3412	1/28/98	463	RTA00000177AR.k.23.4	M00001352D:D02	35550
3412	1/28/98	168	RTA00000177AR.k.23.1	M00001352D:D02	35550
3413	1/28/98	600	RTA00000177AF.g.4.1	M00001346B:B07	4119
3414	1/28/98	280	RTA00000193AF.b.18.1	M00004233C:H09	7542
3415	1/28/98	748	RTA00000198AF.b.24.1	M00001571D:B11	19047
3415	1/28/98	623	RTA00000198R.b.24.1	M00001571D:B11	19047
3416	1/28/98	249	RTA00000200R.o.03.2	M00004257C:H06	22807
3416	1/28/98	178	RTA00000200F.o.03.1	M00004257C:H06	22807
3416	1/28/98	85	RTA00000200R.o.03.1	M00004257C:H06	22807
3417	1/28/98	282	RTA00000193AF.c.15.1	M00004248B:E08	3726
3418	1/28/98	307	RTA00000200F.n.05.2	M00004246C:A09	18989
3418	1/28/98	319	RTA00000200F.n.05.1	M00004246C:A09	18989
3419	1/28/98	319	RTA00000200F.n.05.1	M00004246C:A09	18989

NO: Date of NO: in Priority Priority Appln Appln	ID
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3429 1/28/98 307 RTA00000200F.n.05.2 M00004246C:A09 3429 1/28/98 319 RTA00000200F.n.05.1 M00004246C:A09	18989
110000121001109	18989
3430 1/28/98 52 RTA00000201R.b.02.1 M00004319D:G09	22660
3431 1/28/98 271 RTA00000201F.d.02.1 M00004375A:H01	2599
3431 1/28/98 239 RTA00000201R.d.02.1 M00004375A:H01	2599
3431 1/28/98 227 RTA00000201R.d.02.2 M00004375A:H01 3432 1/28/98 227 RTA00000201R.d.02.2 M00004375A:H01	2599
171000 157571,1101	2599
3432 1/28/98 239 RTA00000201R.d.02.1 M00004375A:H01	2599
3432 1/28/98 271 RTA00000201F.d.02.1 M00004375A:H01	2599
3433 1/28/98 227 RTA00000201R.d.02.2 M00004375A:H01 3433 1/28/98 271 RTA00000201F.d.02.1 M00004375A:H01	2599
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2.10.5	2599
3435 1/28/98 271 RTA00000201F.d.02.1 M00004375A:H01 3435 1/28/98 239 RTA00000201R.d.02.1 M00004375A:H01	2599
3436 1/28/98 178 RTA0000020F.o.03.1 M00004257C:H06	2599
3436 1/28/98 249 RTA00000200R.o.03.2 M00004257C:H06	22807
3436 1/28/98 85 RTA00000200R.o.03.1 M00004257C:H06	22807 22807
3437 1/28/98 273 RTA00000201F.c.08.1 M00004257C:H00	0
3438 1/28/98 328 RTA00000200AF.g.09.1 M00004333C.H07	22785
3438 1/28/98 26 RTA00000200R.g.09.1 M00004131B:H09	22785
3439 2/24/98 571 RTA00000355R.e.14.1 M00004314B:G07	16837
3439 1/28/98 343 RTA00000201F.a.18.1 M00004314B:G07	16837
3440 1/28/98 343 RTA00000201F.a.18.1 M00004314B:G07	16837
3440 2/24/98 571 RTA00000355R.e.14.1 M00004314B:G07	16837
3441 1/28/98 164 RTA00000193AR.i.14.4 M00004307C:A06	9457
3442 1/28/98 50 RTA00000201R.a.02.1 M00004295B:D02	35362
3442 1/28/98 235 RTA00000201AF.a.02.1 M00004295B:D02	35362
3443 1/28/98 235 RTA00000201AF.a.02.1 M00004295B:D02	35362
	22202
3443 1/28/98 50 RTA00000201R.a.02.1 M00004295B:D02	35362

SEQ ID NO:	Filing Date of Priority Appln	SEQ ID NO: in Priority Appln	Sequence Name	Clone Name	Cluster ID
3444	1/28/98	235	RTA00000201AF.a.02.1	M00004295B:D02	35362
3445	1/28/98	227	RTA00000201R.d.02.2	M00004375A:H01	2599
3445	1/28/98	271	RTA00000201F.d.02.1	M00004375A:H01	2599
3445	1/28/98	239	RTA000002011.d.02.1	M00004375A:H01	2599
3446	1/28/98	13	RTA00000190AF.i.5.1	M00003919A:A10	0
3440 3447	1/28/98	72	RTA00000190A1.i.5.1	M000039797X:710	36952
3448	1/28/98	101	RTA000002001.a.d.1 RTA00000191AF.d.08.2	M00004027B:F17	970
3449	1/28/98	79	RTA00000191AI .d.00.2	M00003997B:G07	10282
3450	1/28/98	121	RTA00000199AF.o.16.1	M00003903C:F01	16721
3450 3451	1/28/98	193	RTA00000199AF.n.3.1	M00003979A:103	0
3451	1/28/98	165	RTA00000199AT.II.3.1 RTA00000192AF.g.23.1	M00003340D:C11 M00004157C:A09	6455
3452 3453	1/28/98	381	RTA00000192AI.g.23.1 RTA00000199AF.m.14.1	M00004137C:A07	10580
3453 3454	1/28/98	123	RTA00000199AI.iii.14.1 RTA00000191AF.k.6.1	M00003938A:B04	5451
3454 3455	1/28/98	102	RTA00000191A1.k.o.1 RTA00000199R.j.08.1	M00004078B:A11	37844
	1/28/98	86	RTA00000139KJ.08.1 RTA00000189AF.1.22.1	M00003884D:G07	33333
3456 3457	1/28/98	80 148	RTA00000189AF.1.22.1 RTA00000199F.h.17.2	M00003879C.G10	36254
	1/28/98	143	RTA00000199P.h.17.2 RTA00000199R.h.09.1	M00003877A:A03	76020
3458		143 266	RTA00000199R.ii.09.1 RTA00000199F.f.21.2	M00003807C:H09 M00003847C:E09	13344
3459	1/28/98	153		M00003847C.E09	22561
3460	2/24/98 1/28/98	153 292	RTA00000422F.g.22.1 RTA00000199AF.m.18.1	M00001383B:A06 M00003939C:F04	0
3461		292 275	RTA00000199AF.iii.18.1 RTA00000191AF.o.17.2	M00003939C:F04 M00004102A:H02	5957
3462	1/28/98	273 274			5957 5957
3462	1/28/98	274	RTA00000191AF.o.17.1	M00004102A:H02 M00004375A:H01	2599
3463	1/28/98	239 271	RTA00000201R.d.02.1 RTA00000201F.d.02.1	M00004375A:H01	2599 2599
3463	1/28/98 1/28/98	271	RTA00000201F.d.02.1 RTA00000201R.d.02.2	M00004375A:H01	2599 2599
3463	1/28/98	328	RTA00000201R.d.02.2 RTA00000200AF.g.09.1	M00004373A:H01 M00004131B:H09	2399 22785
3464 3464	1/28/98	328 26	RTA00000200AF.g.09.1 RTA00000200R.g.09.1	M00004131B:H09	22785
3464 3465	1/28/98	214	RTA00000200R.g.09.1 RTA00000200AF.f.22.1	M00004131B:H09	16521
3466	1/28/98	160	RTA00000200AF.1.22.1 RTA00000192AF.b.20.1	M00004121C.100 M00004118D:E08	0
3467	1/28/98	98	RTA00000192AF.6.20.1 RTA00000200AF.f.14.1	M00004118D:C08	22051
3467 3467	1/28/98	100	RTA00000200AF.1.14.1 RTA00000200R.f.14.1	M00004115D:C08	22051
3468	1/28/98	98	RTA00000200K.T.14.1	M00004115D:C08	22051
3468	1/28/98	100	RTA00000200Ar.1.14.1	M00004115D:C08	22051
3469	1/28/98	305	RTA00000200R.1.14.1 RTA00000200AF.b.15.1	M00004113D:C08	10627
3470	1/28/98	98	RTA00000200AF.f.14.1	M00004040D:101 M00004115D:C08	22051
3470	1/28/98	100	RTA00000200AI.I.14.1 RTA00000200R.f.14.1	M00004115D:C08	22051
3471	1/28/98	29	RTA00000200AF.b.19.1	M00004113D:C08	22847
3472	1/28/98	274	RTA00000191AF.o.17.1	M00004042D:H02 M00004102A:H02	5957
3472	1/28/98	275	RTA00000191AF.o.17.2	M00004102A:H02	5957
3473	1/28/98	274	RTA00000191AF.o.17.1	M00004102A:H02	5957
3473	1/28/98	275	RTA00000191AF.o.17.2	M00004102A:H02	5957
3474	1/28/98	275	RTA00000191AF.o.17.2	M00004102A:H02	5957
3474	1/28/98	274	RTA00000191AF.o.17.1	M00004102A:H02	5957
3475	1/28/98	226	RTA00000191AR.o.09.4	M000041027:1102 M00004096A:G02	0
3476	1/28/98	40	RTA00000200AR.e.02.1	M00004090A:F09	36059
3477	1/28/98	175	RTA00000200F.i.5.1	M00004156B:A12	22892
3478	1/28/98	98	RTA00000200AF.f.14.1	M00004115D:C08	22051
3478	1/28/98	100	RTA00000200R.f.14.1	M00004115D:C08	22051
3479	1/28/98	643	RTA00000184AF.c.9.1	M00001546C:G10	16245
3480	1/28/98	615	RTA00000197R.p.20.1	M00001554B:B07	22795
				· · ·	

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln	DT 4 00000107 4 F 20 1	1400001554D.D07	22705
3480	1/28/98	559	RTA00000197AF.p.20.1	M00001554B:B07	22795 6013
3481	1/28/98	660	RTA00000197AF.p.16.1	M00001552D:G08	
3482	1/28/98	521	RTA00000197AF.p.12.1	M00001552B:G05	0
3483	1/28/98	403	RTA00000184AF.f.13.1	M00001550D:H02	3784
3484	1/28/98	517	RTA00000184AF.e.14.1	M00001549C:D02	16347
3485	1/28/98	676	RTA00000197AR.m.14.1	M00001531B:E09	14879
3486	1/28/98	596	RTA00000184AF.d.9.1	M00001548A:B11	6515
3487	1/28/98	559	RTA00000197AF.p.20.1	M00001554B:B07	22795
3487	1/28/98	615	RTA00000197R.p.20.1	M00001554B:B07	22795
3488	1/28/98	729	RTA00000184AF.a.19.1	M00001544C:C06	2628
3489	1/28/98	682	RTA00000125A.j.16.1	M00001544A:E06	0
3490	1/28/98	723	RTA00000183AF.p.24.1	M00001543C:F01	3116
3491	1/28/98	509	RTA00000183AF.p.17.1	M00001543A:H12	5158
3492	1/28/98	738	RTA00000183AF.o.8.1	M00001540C:B10	8927
3493	1/28/98	227	RTA00000201R.d.02.2	M00004375A:H01	2599
3493	1/28/98	271	RTA00000201F.d.02.1	M00004375A:H01	2599
3493	1/28/98	239	RTA00000201R.d.02.1	M00004375A:H01	2599
3494	1/28/98	502	RTA00000197AF.o.23.1	M00001549A:A09	12682
3495	1/28/98	468	RTA00000198AF.a.18.1	M00001561C:E11	0
3496	1/28/98	210	RTA00000189AR.b.12.1	M00003829B:G03	17233
3496	1/28/98	422	RTA00000189AF.b.12.1	M00003829B:G03	17233
3497	1/28/98	748	RTA00000198AF.b.24.1	M00001571D:B11	19047
3497	1/28/98	623	RTA00000198R.b.24.1	M00001571D:B11	19047
3498	1/28/98	397	RTA00000198AF.b.22.1	M00001571B:E03	38956
3499	1/28/98	571	RTA00000198AF.b.14.1	M00001569C:B06	801
3500	1/28/98	492	RTA00000198AF.b.8.1	M00001567C:H12	22636
3500	1/28/98	23	RTA00000198R.b.08.1	M00001567C:H12	22636
3501	1/28/98	492	RTA00000198AF.b.8.1	M00001567C:H12	22636
3501	1/28/98	23	RTA00000198R.b.08.1	M00001567C:H12	22636
3502	1/28/98	559	RTA00000197AF.p.20.1	M00001554B:B07	22795
3502	1/28/98	615	RTA00000197R.p.20.1	M00001554B:B07	22795
3503	1/28/98	727	RTA00000184AF.n.12.2	M00001561D:C11	3727
3504	1/28/98	559	RTA00000197AF.p.20.1	M00001554B:B07	22795
3504	1/28/98	615	RTA00000197R.p.20.1	M00001554B:B07	22795
3505	1/28/98	641	RTA00000198F.a.10.1	M00001558A:E11	6695
3506	1/28/98	731	RTA00000184F.k.02.1	M00001557B:H10	5192
3507	1/28/98	597	RTA00000198F.a.4.1	M00001557A:F01	9635
3508	1/28/98	560	RTA00000184AF.i.23.3	M00001556A:F11	1577
3509	1/28/98	60 1	RTA00000184AF.i.10.2	M00001555A:B01	3744
3510	1/28/98	700	RTA00000183AF.i.18.2	M00001529D:H02	40129
3511	1/28/98	437	RTA00000198R.a.23.1	M00001563B:D11	10700
3512	1/28/98	591	RTA00000197AF.h.1.1	M00001470A:H01	13075
3512	1/28/98	110	RTA00000197R.h.01.1	M00001470A:H01	13075
3513	1/28/98	259	RTA00000197AF.j.4.1	M00001492D:A11	17209
3513	1/28/98	386	RTA00000197AR.j.04.1	M00001492D:A11	17209
3514	1/28/98	386	RTA00000197AR.j.04.1	M00001492D:A11	17209
3514	1/28/98	259	RTA00000197AF.j.4.1	M00001492D:A11	17209
3515	1/28/98	644	RTA00000197F.i.12.1	M00001489B:A06	3605
3516	1/28/98	633	RTA00000197F.i.8.1	M00001488A:E01	6292
3517	1/28/98	546	RTA00000197F.i.6.1	M00001487C:D06	12149

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
2510	Appln 1/28/98	Appln 650	RTA00000183AR.n.17.1	M00001539B:H06	9800
3518 3519	1/28/98	513	RTA00000197AF.h.14.1	M00001337B:F04	7045
3520	1/28/98	519	RTA0000013771	M00001499B:A11	10539
3520 3521	1/28/98	110	RTA00000197R.h.01.1	M00001470A:H01	13075
3521	1/28/98	591	RTA00000197AF.h.1.1	M00001470A:H01	13075
	1/28/98	446	RTA00000137AF.ii.111 RTA00000182AF.a.23.3	M00001463A:F06	9755
3522	1/28/98	739	RTA00000182AF.a.25.3 RTA00000181AF.p.12.3	M000014697t:100	22204
3523	1/28/98	635	RTA00000181AF.p.7.3	M00001460A:E01	38773
3524	1/28/98	720	RTA00000197AF.f.14.1	M00001459B:C09	3732
3525		623	RTA00000197A1.1.14.1 RTA00000198R.b.24.1	M00001437D:B11	19047
3526	1/28/98 1/28/98	748	RTA00000198AF.b.24.1	M00001571D:B11	19047
3526	1/28/98	419	RTA00000138AF.j.20.1	M00001371B:D11	4769
3527	1/28/98	632	RTA00000182A1.j.20.1 RTA00000183AR.g.03.1	M00001403D:D03	3956
3528		630	RTA00000183AR.g.03.1	M00001512D:G09	3956
3528	1/28/98	695	RTA00000183AR.g.03.2 RTA00000197F.m.5.1	M00001512D:G09	10872
3529	1/28/98	693 479	RTA00000197F.II.3.1 RTA00000197R.I.22.1	M00001528A:C11	6962
3530	1/28/98			M00001528A:C11	6962
3530	1/28/98	665	RTA00000197AF.l.22.1	M00001528A:C11	6962
3531	1/28/98	479	RTA00000197R.1.22.1 RTA00000197AF.1.22.1	M00001528A:C11	6962
3531	1/28/98	665	RTA00000197AF.I.22.1 RTA00000197R.I.22.1	M00001528A:C11	6962
3532	1/28/98	479		M00001528A:C11	6962
3532	1/28/98	665	RTA00000197AF.l.22.1	M00001528A:C11	6962
3533	1/28/98	479	RTA00000197R.l.22.1 RTA00000197AF.l.22.1	M00001528A:C11	6962
3533	1/28/98	665 550	RTA00000197AF.1.22.1 RTA00000183AF.g.14.1	M00001528A.C11 M00001513D:A03	0
3534	1/28/98		RTA00000195AF.b.6.1	M00001313D:A03	39490
3535	1/28/98	404	RTA00000193AF.b.6.1 RTA00000183AR.g.03.2	M00001490C.G10 M00001512D:G09	3956
3536	1/28/98	630 632	RTA00000183AR.g.03.1	M00001512D:G09	3956
3536	1/28/98	570	RTA00000183AF.a.19.2	M00001312D:G09 M00001499A:A05	3788
3537	1/28/98	630	RTA00000183AR.g.03.2	M00001439A:A03	3956
3538	1/28/98 1/28/98	632	RTA00000183AR.g.03.1	M00001512D:G09	3956
3538 3539	1/28/98	603	RTA00000183AR.d.11.3	M00001512D:G09	6420
3539 3540	1/28/98	715	RTA00000197AR.k.11.1	M00001504D:G00 M00001500D:E10	53758
3540 3541	1/28/98	503	RTA00000197AR.k.11.1	M00001500D:D10	3138
3542	1/28/98	719	RTA0000013771 .k.511	M00001500A:B02	0
3542 3543	1/28/98	271	RTA00000103A1:0:12:1	M00004375A:H01	2599
3543	1/28/98	239	RTA00000201R.d.02.1	M00004375A:H01	2599
3543	1/28/98	227	RTA00000201R.d.02.2	M00004375A:H01	2599
3544	1/28/98	630	RTA00000183AR.g.03.2	M00001512D:G09	3956
3544	1/28/98	632	RTA00000183AR.g.03.1	M00001512D:G09	3956
3545	3/24/98	15	RTA00000425F.j.14.1	M00001639D:C12	73397
3546	3/24/98	111	RTA00000425F.d.08.1	M00001631A:F06	74350
3547	3/24/98	152	RTA00000425F.d.07.1	M00001631A:F12	43197
3548	3/24/98	147	RTA00000425F.d.21.1	M00001631B:H04	78920
3549	3/24/98	77	RTA00000425F.i.17.1	M00001633A:F11	43213
3550	3/24/98	418	RTA00000425F.i.18.1	M00001633A:G10	42255
3551	3/24/98	197	RTA00000425F.j.20.1	M00001633B:A12	26760
3552	3/24/98	143	RTA00000425F.j.22.1	M00001633B:E03	73882
3553	3/24/98	283	RTA00000425F.k.20.1	M00001633C:A08	74048
3554	3/24/98	139	RTA00000425F.k.22.1	M00001633C:E12	78123
3555	2/24/98	870	RTA00000418F.n.24.1	M00001659D:C09	73153

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
222	Appln	Appln	D.T. 4.000.004.0555 1.5.1	1.000001.00C1 TI10	70004
3556	3/24/98	403	RTA00000425F.n.17.1	M00001636A:H12	78304
3557	2/24/98	1109	RTA00000422F.f.22.1	M00001584A:G03	38703
3558	3/24/98	150	RTA00000424F.d.04.1	M00001478A:F12	76505
3558	3/24/98	149	RTA00000424F.d.04.3	M00001478A:F12	76505
3559	3/24/98	358	RTA00000425F.n.19.1	M00001638B:C08	78324
3560	3/24/98	165	RTA00000425F.e.21.1	M00001629D:D10	77203
3561	3/24/98	443	RTA00000425F.k.16.1	M00001640A:F05	75282
3562	3/24/98	252	RTA00000425F.m.03.1	M00001642D:G08	76045
3563	3/24/98	116	RTA00000425F.n.05.1	M00001647D:G07	73965
3564	3/24/98	425	RTA00000522F.i.07.2	M00001649A:E10	78377
3565	3/24/98	445	RTA00000522F.j.08.2	M00001650D:D10	76613
3566	3/24/98	371	RTA00000522F.j.09.2	M00001650D:F11	78522
3567	3/24/98	97	RTA00000522F.j.14.2	M00001651C:D11	73123
3568	3/24/98	69	RTA00000522F.j.15.2	M00001651C:G12	76535
3569	3/24/98	373	RTA00000522F.j.19.2	M00001652B:D06	76224
3570	3/24/98	93	RTA00000522F.k.14.1	M00001652D:G02	74280
3571	3/24/98	50	RTA00000522F.k.15.1	M00001652D:G06	76866
3572	3/24/98	141	RTA00000522F.k.19.1	M00001653A:A05	32625
3573	3/24/98	409	RTA00000425F.I.10.1	M00001638A:C08	26893
3574	2/24/98	443	RTA00000414F.f.15.1	M00005260A:A12	0
3575	2/24/98	886	RTA00000420F.m.15.1	M00005235B:F10	0
3576	2/24/98	260	RTA00000414F.e.08.1	M00005236A:E04	Ö
3577	2/24/98	734	RTA00000414F.e.09.1	M00005236A:G10	Ŏ
3578	2/24/98	1077	RTA00000414F.e.11.1	M00005236B:A12	Ö
3579	2/24/98	970	RTA00000414F.e.15.1	M00005236B:G03	Ŏ
3580	2/24/98	271	RTA00000414F.e.16.1	M00005236B:H10	0
3581	2/24/98	58	RTA00000420F.m.18.1	M00005254D:A10	Ö
3582	2/24/98	289	RTA00000420F.n.08.1	M00005257A:H11	0
3583	2/24/98	1033	RTA00000414F.e.19.1	M00005257C:E05	Ŏ
3584	2/24/98	793	RTA00000414F.e.21.1	M00005257C:G01	0
3585	2/24/98	36	RTA00000414F.e.22.1	M00005257D:A06	Ŏ
3586	2/24/98	852	RTA00000414F.f.03.1	M00005257D:700 M00005257D:G07	0
3587	3/24/98	341	RTA000004141.1.03.1 RTA00000425F.d.06.1	M00003237B:G07	77660
3588	2/24/98	961	RTA00000420F.n.21.2	M00005259B:D12	0
3589	3/24/98	441	RTA00000420F.m.21.2 RTA00000528F.g.22.2	M00003237B:B12	920
3590	2/24/98	940	RTA000003201.g.22.2	M000070502:109	0
3591	2/24/98	160	RTA00000414F.f.19.1	M00005260B:E11	0
3592	3/24/98	140	RTA000004141.1.19.11 RTA00000424F.m.14.1	M00003200B:E11	77491
3593	3/24/98	34	RTA00000424F.m.15.1	M00001612D:F06	73759
3594	3/24/98	212	RTA00000424F.n.06.1	M00001612B:100	74737
3595	3/24/98	308	RTA00000424F.k.23.1	M00001613A:B02	31061
3596	3/24/98	372	RTA00000424F.m.24.1	M00001614A:B10	77045
3597	3/24/98	396	RTA00000524F.m.24.1 RTA00000528F.g.05.2	M00001614C:G07	3770
3598	3/24/98	296	RTA000003261.g.03.2 RTA00000425F.e.02.1	M00001615C:E07	76143
3599	3/24/98	99	RTA00000425F.c.20.1	M00001625C.F10 M00001626D:A02	73581
3600	3/24/98	442	RTA00000425F.d.14.1	M00001629D:A02	13417
3601	3/24/98	357	RTA00000425F.d.14.1 RTA00000425F.e.19.1	M00001629A:H09 M00001629D:B10	73409
3602	2/24/98	210	RTA00000425F.e.19.1 RTA00000419F.p.24.1	M00001029D:B10 M00004039B:E12	63477
3603	2/24/98	501	RTA00000419F.p.24.1 RTA00000414F.f.05.1	M00004039B:E12 M00005257D:H11	034//
3604	2/24/98	561	RTA00000414F.I.03.1 RTA00000420F.e.10.1	M00003237D:H11 M00004108D:G04	65899
JUU 1	4147170	501	K1/A000007201.6.10.1	14100001 F000014	ひょりブブ

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
3605	2/24/98	758	RTA00000407F.a.01.1	M00004039A:H11	12501
3606	2/24/98	688	RTA00000413F.d.23.1	M00004090B:H06	66030
3607	2/24/98	124	RTA00000420F.d.05.1	M00004092B:E05	64432
3608	2/24/98	329	RTA00000413F.e.16.1	M00004093C:C02	63836
3609	2/24/98	359	RTA00000420F.d.12.1	M00004096D:H03	64095
3610	2/24/98	429	RTA00000422F.c.17.1	M00004099D:F01	1360
3611	2/24/98	630	RTA00000413F.f.19.1	M00004100B:C07	65189
3612	2/24/98	439	RTA00000413F.g.23.1	M00004103B:E09	40700
3613	2/24/98	3	RTA00000420F.d.18.1	M00004105C:B05	63074
3614	2/24/98	1064	RTA00000420F.d.19.1	M00004105C:C08	43146
3615	2/24/98	671	RTA00000413F.h.12.1	M00004107A:A12	66929
3616	2/24/98	507	RTA00000420F.e.02.1	M00004107B:D07	40259
3617	2/24/98	319	RTA00000420F.b.21.1	M00004088D:B10	65057
3618	2/24/98	931	RTA00000420F.e.09.1	M00004108D:E07	66325
3619	2/24/98	840	RTA00000420F.b.20.1	M00004088D:B05	0
3620	2/24/98	545	RTA00000420F.e.15.1	M00004110A:A10	20190
3621	2/24/98	981	RTA00000420F.e.20.1	M00004110B:A07	64762
3622	3/24/98	370	RTA00000424F.d.19.3	M00001448B:A07	73180
3623	3/24/98	370	RTA00000424F.d.19.3	M00001448B:A07	73180
3624	3/24/98	189	RTA00000424F.d.22.3	M00001448B:G07	76189
3625	3/24/98	189	RTA00000424F.d.22.3	M00001448B:G07	76189
3626	3/24/98	92	RTA00000424F.a.24.4	M00001448D:E11	73951
3627	3/24/98	92	RTA00000424F.a.24.4	M00001448D:E11	73951
3628	3/24/98	279	RTA00000528F.b.03.1	M00001455A:D10	2078
3629	3/24/98	279	RTA00000528F.b.03.1	M00001455A:D10	2078
3630	3/24/98	480	RTA00000424F.d.17.3	M00001455A:E11	73958
3631	3/24/98	480	RTA00000424F.d.17.3	M00001455A:E11	73958
3632	2/24/98	583	RTA00000406F.i.17.1	M00003904B:C03	37902
3633	2/24/98	590	RTA00000407F.b.22.1	M00004108B:B02	37487
3634	2/24/98	1075	RTA00000413F.b.17.1	M00004078A:F07	21704
3635	2/24/98	544	RTA00000420F.1.21.2	M00005232A:H12	0
3636	1/28/98	684	RTA00000200AR.b.11.1	M00004040A:G12	12043
3636	2/24/98	1166	RTA00000347F.h.01.1	M00004040A:G12	12043
3637	1/28/98	684	RTA00000200AR.b.11.1	M00004040A:G12	12043
3637	2/24/98	1166	RTA00000347F.h.01.1	M00004040A:G12	12043
3638	2/24/98	1087	RTA00000401F.o.13.1	M00004040C:A01	3220
3639	2/24/98	114	RTA00000341F.m.21.1	M00004051D:E01	0
3640	2/24/98	811	RTA00000413F.a.12.1	M00004072D:F09	63403
3641	2/24/98	714	RTA00000420F.a.08.1	M00004073A:D10	19473
3642	1/28/98	387	RTA00000191AF.j.14.1	M00004073A:H12	1002
3642	2/24/98	632	RTA00000191AF.j.14.1	M00004073A:H12	1002
3643	1/28/98	387	RTA00000191AF.j.14.1	M00004073A:H12	1002
3643	2/24/98	632	RTA00000191AF.j.14.1	M00004073A:H12	1002
3644	2/24/98	964	RTA00000423F.I.15.1	M00004075B:G09	11219
3645	2/24/98	355	RTA00000420F.a.19.1	M00004076A:D12	34192
3646	2/24/98	745	RTA00000413F.b.04.1	M00004076D:H07	66427
3647	2/24/98	64	RTA00000413F.d.18.1	M00004090B:B04	65305
3648	2/24/98	698	RTA00000413F.b.16.1	M00004078A:E05	65126
3649	2/24/98	190	RTA00000419F.p.23.1	M00004039B:A05	64748
3650	2/24/98	903	RTA00000420F.a.21.1	M00004078B:C11	66241

SEQ ID	Filing	SEQ ID	Sequence Name	Clone Name	Cluster
NO:	Date of	NO: in			ID
	Priority	Priority			
2651	Appln 2/24/98	Appln 588	PTA00000420E - 22 1	M00004070D.E13	40160
3651 3652	2/24/98	588 1185	RTA00000420F.a.23.1 RTA00000413F.b.20.1	M00004078B:F12 M00004079D:G08	42158 66063
3652 3653	2/24/98	619	RTA00000413F.b.20.1 RTA00000420F.b.04.1	M00004079D:G08 M00004081A:E02	63820
		988			15570
3654	2/24/98		RTA00000407F.a.22.1 RTA00000407F.a.23.1	M00004081A:G01	
3655	2/24/98	705		M00004081C:A10	23489
3656	2/24/98	282	RTA00000407F.a.24.1	M00004083A:E08	37560
3657	2/24/98	835	RTA00000413F.c.10.1	M00004083B:C01	65600
3658	2/24/98	598	RTA00000420F.b.18.1	M00004086D:G08	66136
3659	2/24/98	335	RTA00000413F.d.02.1	M00004087B:A12	66172
3660	2/24/98	504	RTA00000413F.d.05.1	M00004087C:A01	64788
3661	2/24/98	76	RTA00000413F.d.16.1	M00004088C:F01	63331
3662	2/24/98	726	RTA00000420F.b.19.1	M00004088D:A11	36873
3663	2/24/98	521	RTA00000413F.b.14.1	M00004078A:C11	66591
3664	2/24/98	255	RTA00000419F.o.16.1	M00003989C:G05	62867
3665	2/24/98	665	RTA00000419F.p.20.1	M00004039A:C03	9458
3666	2/24/98	1234	RTA00000352R.c.20.1	M00003982A:B12	7339
3667	2/24/98	247	RTA00000412F.j.17.1	M00003982C:G04	64071
3668	2/24/98	1145	RTA00000423F.k.21.2	M00003984D:B08	37499
3669	2/24/98	993	RTA00000406F.o.05.1	M00003985B:G04	37894
3670	2/24/98	328	RTA00000423F.k.19.2	M00003985D:E10	17615
3671	2/24/98	254	RTA00000341F.1.15.1	M00003986B:A08	5294
3672	2/24/98	948	RTA00000419F.o.06.1	M00003986C:D09	64643
3673	2/24/98	661	RTA00000341F.I.16.1	M00003986D:C08	8479
3674	2/24/98	117	RTA00000341F.m.13.1	M00003987B:E12	26502
3675	2/24/98	1210	RTA00000419F.o.09.1	M00003987B:F08	66396
3676	2/24/98	460	RTA00000341F.j.12.1	M00003987C:G03	12195
3677	2/24/98	486	RTA00000346F.I.13.1	M00003980B:C11	7542
3678	2/24/98	723	RTA00000419F.o.15.1	M00003989C:D03	32487
3679	2/24/98	897	RTA00000419F.n.24.1	M00003980A:F04	65995
3680	2/24/98	92	RTA00000412F.1.04.1	M00003989D:F12	66372
3681	2/24/98	1014	RTA00000412F.1.14.1	M00004029B:F01	62792
3682	2/24/98	348	RTA00000412F.I.19.1	M00004029C:C05	65825
3683	2/24/98	284	RTA00000412F.1.21.1	M00004029C:G10	65183
3684	2/24/98	188	RTA00000406F.p.04.1	M00004030D:F11	37458
3685	2/24/98	812	RTA00000412F.o.05.1	M00004034A:A01	63575
3686	2/24/98	911	RTA00000406F.p.13.1	M00004034C:G02	8584
3687	2/24/98	230	RTA00000423F.k.01.1	M00004034D:E09	40426
3688	2/24/98	1076	RTA00000423F.k.09.1	M00004035B:H09	26630
3689	2/24/98	941	RTA00000419F.p.08.1	M00004036D:B04	65560
3690	2/24/98	1186	RTA00000419F.p.10.1	M00004036D:B09	41448
3691	2/24/98	42	RTA00000423F.k.17.2	M00004038A:F02	37512
3692	2/24/98	934	RTA00000414F.e.01.1	M00005233D:H07	0
3693	2/24/98	37	RTA00000406F.o.15.1	M00003988D:A08	37482
3694	2/24/98	1016	RTA00000406F.n.12.1	M00003960A:G07	37517
3695	1/28/98	584	RTA00000190AR.c.03.1	M00003904C:A08	0
3695	2/24/98	1069	RTA00000346F.k.05.1	M00003904C:A08	0
3696	1/28/98	584	RTA00000190AR.c.03.1	M00003904C:A08	0
3696	2/24/98	1069	RTA00000346F.k.05.1	M00003904C:A08	0
3697	2/24/98	489	RTA00000406F.j.19.1	M00003906A:F12	1685
3698	2/24/98	461	RTA00000412F.d.16.1	M00003906B:H06	26829

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
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3699	2/24/98	558	RTA00000419F.m.04.1	M00003906C:C05	74367
3700	2/24/98	120	RTA00000401F.m.02.1	M00003907A:F01	1573
3701	2/24/98	628	RTA00000412F.d.19.1	M00003907B:C03	75743
3702	2/24/98	792	RTA00000406F.k.11.1	M00003907B:D05	38715
3703	2/24/98	292	RTA00000423F.i.18.1	M00003918A:D08	14996
3704	2/24/98	1192	RTA00000406F.m.17.1	M00003918A:F09	0
3705	2/24/98	9	RTA00000406F.n.02.1	M00003918C:H10	15051
3706	2/24/98	629	RTA00000352R.c.04.1	M00003924A:D08	71976
3707	2/24/98	438	RTA00000195R.d.09.1	M00003981C:B04	8537
3708	2/24/98	433	RTA00000419F.n.02.1	M00003958B:H08	65963
3709	2/24/98	147	RTA00000422F.c.02.1	M00004118B:A03	2902
3710	2/24/98	649	RTA00000412F.g.03.1	M00003971B:A10	64740
3711	2/24/98	1141	RTA00000347F.f.08.1	M00003972D:H02	5948
3712	2/24/98	252	RTA00000412F.g.24.1	M00003973C:C03	28741
3713	2/24/98	732	RTA00000412F.h.11.1	M00003974B:B11	63175
3714	2/24/98	181	RTA00000412F.h.21.1	M00003974D:F02	64348
3715	2/24/98	345	RTA00000412F.h.23.2	M00003974D:H04	65118
3716	2/24/98	148	RTA00000419F.n.04.1	M00003975C:F07	13102
3717	2/24/98	311	RTA00000419F.n.09.1	M00003977C:A06	66070
3718	2/24/98	1044	RTA00000419F.n.11.1	M00003977C:B03	66477
3719	2/24/98	652	RTA00000419F.n.12.1	M00003977D:A03	66086
3720	2/24/98	452	RTA00000419F.n.13.1	M00003977D:A06	66026
3721	2/24/98	796	RTA00000419F.n.15.1	M00003977D:D04	63484
3722	2/24/98	1249	RTA00000419F.n.17.1	M00003978D:G04	63186
3723	2/24/98	860	RTA00000419F.m.23.1	M00003958B:E11	64263
3724	2/24/98	713	RTA00000414F.b.10.1	M00005212D:D09	0
3725	2/24/98	1002	RTA00000419F.p.18.1	M00004038D:G06	63002
3726	2/24/98	1048	RTA00000413F.o.07.2	M00005100A:C01	0
3727	2/24/98	508	RTA00000420F.i.20.1	M00005101C:E12	0
3728	2/24/98	1001	RTA00000413F.p.07.2	M00005102C:D03	0
3729	2/24/98	88	RTA00000420F.i.24.1	M00005134B:E08	0
3730	2/24/98	93	RTA00000413F.p.24.1	M00005139A:H03	0
3731	2/24/98	142	RTA00000420F.j.19.1	M00005140C:B10	0
3732	2/24/98	833	RTA00000420F.j.20.1	M00005140D:C06	0
3733	2/24/98	316	RTA00000414F.a.02.1	M00005178D:H04	0
3734	2/24/98	1100	RTA00000414F.a.12.1	M00005210A:E06	0
3735	2/24/98	1175	RTA00000414F.b.04.1	M00005212B:E01	0
3736	2/24/98	1236	RTA00000414F.b.06.1	M00005212C:C03	0
3737	2/24/98	747	RTA00000413F.n.24.1	M00004960C:E10	0
3738	2/24/98	207	RTA00000414F.b.08.1	M00005212C:H02	0
3739	2/24/98	935	RTA00000420F.i.07.1	M00004960A:B08	0
3740	2/24/98	741	RTA00000414F.b.12.1	M00005212D:H01	0
3741	2/24/98	865	RTA00000414F.c.03.1	M00005216A:D09	0
3742	2/24/98	862	RTA00000414F.c.07.1	M00005216A:H01	0
3743	2/24/98	565	RTA00000420F.k.17.2	M00005217B:A06	0
3744	2/24/98	1226	RTA00000414F.c.12.1	M00005218A:F09	0
3745	2/24/98	512	RTA00000414F.c.16.1	M00005228A:B03	0
3746	2/24/98	817	RTA00000420F.1.08.2	M00005228C:C05	0
3747	2/24/98	573	RTA00000414F.c.23.1	M00005229B:G12	0
3748	2/24/98	1237	RTA00000414F.c.24.1	M00005229B:H04	0

Appln	SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
3749 2/24/98 727 RTA00000414F.d.02.1 M00005229B:H06 0 3750 2/24/98 566 RTA00000420F.l.12.2 M00005229B:H09 0 3751 2/24/98 307 RTA00000420F.l.12.2 M00005239B:H09 0 3752 3/24/98 150 RTA00000424F.d.04.1 M00001478A:F12 76505 3753 2/24/98 946 RTA00000424F.d.04.1 M00001478A:F12 76505 3754 1/28/98 343 RTA0000025E.e.14.1 M0000414B:G07 16837 3754 2/24/98 571 RTA0000035E.e.14.1 M0000414B:G07 16837 3755 2/24/98 824 RTA0000040Fr.c.08.1 M00004118B:F01 63073 3757 2/24/98 824 RTA00000346F.n.08.1 M000041199:C09 66312 3759 2/24/98 813 RTA00000346F.n.06.1 M00004139C:A12 12439 3761 2/24/98 368 RTA0000034Fr.n.06.1 M00004139C:A12 12439 3762 1/28/98 685 RT		Priority	Priority			
3750 2/24/98 366	37/10			RTA00000414F 4 02 1	M00005220R·H06	0
3751 2/24/98 307 RTA00000420F.I.12.2 M000005230B:H09 0 3752 3/24/98 149 RTA00000424F.d.04.3 M00001478A:F12 76505 3753 3/24/98 946 RTA00000414F.b.07.1 M00001478A:F12 76505 3754 1/28/98 343 RTA0000021F.a.18.1 M00004314B:G07 16837 3754 2/24/98 571 RTA0000035SR.e.14.1 M00004314B:G07 16837 3755 2/24/98 481 RTA0000047F.c.08.1 M00004118B:F01 63073 3756 2/24/98 841 RTA0000047F.c.08.1 M00004118B:F01 63073 3756 2/24/98 813 RTA00000346F.a.06.1 M00004137A:D06 0 3761 2/24/98 813 RTA00000346F.a.02.1 M00004137A:D06 0 3760 2/24/98 868 RTA00000346F.a.06.1 M00004139C:A12 12439 3761 2/24/98 868 RTA0000035SR.a.12.1 M00004159C:F09 36756 3762 1/28/98 685			. — .			
3752 3/24/98 149 RTA00000424F.d.04.3 M00001478A:F12 76505 3752 3/24/98 150 RTA00000444F.d.04.1 M00001478A:F12 76505 3753 3/24/98 946 RTA00000414F.b.07.1 M00005212C:D02 0 3754 1/28/98 343 RTA00000201F.a.18.1 M00004314B:G07 16837 3754 2/24/98 481 RTA00000355R.e.14.1 M00004314B:G07 16837 3755 2/24/98 481 RTA00000415F.i.23.1 M00004118B:B05 3754 3757 2/24/98 813 RTA00000420F.f.07.1 M00004118D:B05 3754 3757 2/24/98 813 RTA00000346F.a.06.1 M00004136D:B02 4937 3759 2/24/98 1070 RTA00000346F.a.06.1 M00004137A:D06 0 3760 2/24/98 368 RTA00000346F.a.08.1 M00004139C:B02 0 3761 2/24/98 685 RTA0000034F.a.08.1 M00004159C:F09 36756 3762 1/28/98 685 R						
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3753 2/24/98 946 RTA0000021F.a.18.1 M00005212C.D02 0 3754 1/28/98 343 RTA00000201F.a.18.1 M00004314B:G07 16837 3754 2/24/98 571 RTA000000355R.e.14.1 M00004118B:F01 6387 3755 2/24/98 181 RTA00000417F.c.08.1 M00004118B:F01 63073 3757 2/24/98 824 RTA00000420F.f.07.1 M00004118B:F01 66312 3758 2/24/98 813 RTA00000346F.o.06.1 M00004136D:B02 4937 3760 2/24/98 813 RTA00000346F.o.06.1 M00004137A:D06 0 3761 2/24/98 368 RTA00000346F.o.08.1 M00004159C:B02 0 3762 1/28/98 685 RTA00000346F.o.08.1 M00004159C:B02 36756 3763 1/28/98 685 RTA0000035Fs.a.12.1 M00004159C:F09 36756 3763 1/28/98 685 RTA0000035Fs.a.12.1 M00004159C:F09 36756 3764 2/24/98 125						
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3754 2/24/98 571						-
3755 2/24/98 1039 RTA00000413F.i.23.1 M00004118B:F01 63073 3756 2/24/98 1039 RTA00000407F.c.08.1 M00004118D:B05 37549 3757 2/24/98 813 RTA00000340F.c.06.1 M00004118D:B02 4937 3759 2/24/98 813 RTA00000346F.c.06.1 M00004137A:D06 0 0 0 0 0 0 0 0 0						
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3757 2/24/98 824 RTA00000420F.f.07.1 M00004119A:C09 66312 3758 2/24/98 813 RTA00000346F.n.06.1 M00004136D:B02 4937 3759 2/24/98 283 RTA00000346F.n.06.1 M00004139C:A12 12439 3761 2/24/98 368 RTA00000346F.n.08.1 M00004149C:B02 0 3762 2/24/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 3763 1/28/98 685 RTA00000200F.i.9.1 M00004159C:F09 36756 3763 1/28/98 685 RTA0000035FR.a.12.1 M00004159C:F09 36756 3763 1/24/98 1254 RTA0000035FR.a.12.1 M00004159C:F09 36756 3765 2/24/98 1188 RTA0000034IF.p.I1.1 M00004159C:F09 36756 3766 2/24/98 128 RTA0000035ER.a.10.6.1 M00004159C:G12 0 3767 2/24/98 456 RTA0000035ER.a.10.6.1 M00004169D:B11 37189 3768 2/24/98 882						
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3759 2/24/98 1070						
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3777 2/24/98 1238 RTA00000420F.g.09.1 M00004895B:E12 0 3778 2/24/98 18 RTA00000420F.g.12.1 M00004895B:G04 0 3779 2/24/98 1196 RTA00000413F.I.18.1 M00004895D:G07 0 3780 2/24/98 579 RTA00000413F.m.16.1 M00004899D:G06 0 3781 2/24/98 143 RTA00000420F.h.13.1 M00004899D:G06 0 3782 2/24/98 909 RTA00000420F.i.04.1 M00004959D:H12 0 3783 2/24/98 709 RTA00000352R.p.09.1 M00004228C:H03 16915 3784 3/24/98 221 RTA00000427F.j.22.1 M00004097D:B05 66367 3785 3/24/98 188 RTA00000525F.c.15.1 M00004040A:A07 7692 3786 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 3787 3/24/98 325 RTA00000525F.c.17.1 M00004040B:C05 24208 3789 3/24/98 159 RTA0000052						
3778 2/24/98 18 RTA00000420F.g.12.1 M00004895B:G04 0 3779 2/24/98 1196 RTA00000413F.I.18.1 M00004895D:G07 0 3780 2/24/98 579 RTA00000413F.m.16.1 M00004898C:F03 0 3781 2/24/98 143 RTA00000420F.h.13.1 M00004899D:G06 0 3782 2/24/98 909 RTA00000420F.i.04.1 M00004959D:H12 0 3783 2/24/98 709 RTA00000352R.p.09.1 M00004228C:H03 16915 3784 3/24/98 221 RTA00000427F.j.22.1 M00004097D:B05 66367 3785 3/24/98 188 RTA00000525F.c.15.1 M00004040A:A07 7692 3786 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 3787 3/24/98 53 RTA00000525F.c.17.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000					M00004895B:E12	
3779 2/24/98 1196 RTA00000413F.I.18.1 M00004895D:G07 0 3780 2/24/98 579 RTA00000413F.m.16.1 M00004898C:F03 0 3781 2/24/98 143 RTA00000420F.h.13.1 M00004899D:G06 0 3782 2/24/98 909 RTA00000420F.i.04.1 M00004959D:H12 0 3783 2/24/98 709 RTA00000352R.p.09.1 M00004228C:H03 16915 3784 3/24/98 221 RTA00000427F.j.22.1 M00004097D:B05 66367 3785 3/24/98 188 RTA00000525F.c.15.1 M00004040A:A07 7692 3786 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 3787 3/24/98 53 RTA00000525F.c.17.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011		2/24/98	18			
3780 2/24/98 579 RTA00000413F.m.16.1 M00004898C:F03 0 3781 2/24/98 143 RTA00000420F.h.13.1 M00004899D:G06 0 3782 2/24/98 909 RTA00000420F.i.04.1 M00004959D:H12 0 3783 2/24/98 709 RTA00000352R.p.09.1 M00004228C:H03 16915 3784 3/24/98 221 RTA00000427F.j.22.1 M00004097D:B05 66367 3785 3/24/98 188 RTA00000525F.c.15.1 M00004040A:A07 7692 3786 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 3787 3/24/98 53 RTA00000525F.c.17.1 M00004040A:C08 38160 3788 3/24/98 325 RTA00000525F.c.18.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3779	2/24/98	1196		M00004895D:G07	
3782 2/24/98 909 RTA00000420F.i.04.1 M00004959D:H12 0 3783 2/24/98 709 RTA00000352R.p.09.1 M00004228C:H03 16915 3784 3/24/98 221 RTA00000427F.j.22.1 M00004097D:B05 66367 3785 3/24/98 188 RTA00000525F.c.15.1 M00004040A:A07 7692 3786 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 3787 3/24/98 53 RTA00000525F.c.17.1 M00004040A:C08 38160 3788 3/24/98 325 RTA00000525F.c.18.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3780	2/24/98	579		M00004898C:F03	
3783 2/24/98 709 RTA00000352R.p.09.1 M00004228C:H03 16915 3784 3/24/98 221 RTA00000427F.j.22.1 M00004097D:B05 66367 3785 3/24/98 188 RTA00000525F.c.15.1 M00004040A:A07 7692 3786 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 3787 3/24/98 53 RTA00000525F.c.17.1 M00004040A:C08 38160 3788 3/24/98 325 RTA00000525F.c.18.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3781	2/24/98	143	RTA00000420F.h.13.1	M00004899D:G06	0
3784 3/24/98 221 RTA00000427F.j.22.1 M00004097D:B05 66367 3785 3/24/98 188 RTA00000525F.c.15.1 M00004040A:A07 7692 3786 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 3787 3/24/98 53 RTA00000525F.c.17.1 M00004040A:C08 38160 3788 3/24/98 325 RTA00000525F.c.18.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3782	2/24/98	909	RTA00000420F.i.04.1	M00004959D:H12	0
3785 3/24/98 188 RTA00000525F.c.15.1 M00004040A:A07 7692 3786 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 3787 3/24/98 53 RTA00000525F.c.17.1 M00004040A:C08 38160 3788 3/24/98 325 RTA00000525F.c.18.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3783	2/24/98	709	RTA00000352R.p.09.1	M00004228C:H03	16915
3786 3/24/98 401 RTA00000525F.c.16.1 M00004040A:B04 38209 3787 3/24/98 53 RTA00000525F.c.17.1 M00004040A:C08 38160 3788 3/24/98 325 RTA00000525F.c.18.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3784	3/24/98	221	RTA00000427F.j.22.1	M00004097D:B05	66367
3787 3/24/98 53 RTA00000525F.c.17.1 M00004040A:C08 38160 3788 3/24/98 325 RTA00000525F.c.18.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3785	3/24/98	188	RTA00000525F.c.15.1	M00004040A:A07	7692
3788 3/24/98 325 RTA00000525F.c.18.1 M00004040B:C05 24208 3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3786	3/24/98	401	RTA00000525F.c.16.1	M00004040A:B04	38209
3789 3/24/98 159 RTA00000525F.c.19.1 M00004040B:F07 38159 3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3787	3/24/98	53	RTA00000525F.c.17.1	M00004040A:C08	38160
3790 3/24/98 209 RTA00000427F.g.16.1 M00004069A:E12 63011	3788	3/24/98	325	RTA00000525F.c.18.1	M00004040B:C05	24208
	3789	3/24/98	159	RTA00000525F.c.19.1		38159
3791 3/24/98 123 RTA00000427F.g.05.1 M00004069C:C08 63138		3/24/98			M00004069A:E12	
	3791	3/24/98	123	RTA00000427F.g.05.1	M00004069C:C08	63138

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
2702	Appln	Appln	DT 4 00000427F : 10 1	N400004077 A -C12	41206
3792	3/24/98	62	RTA00000427F.j.19.1	M00004077A:G12	41395
3793	3/24/98	265	RTA00000427F.h.02.1	M00004085B:G01	63652
3794	3/24/98	235	RTA00000427F.g.19.1	M00004087A:B05	64611
3795	3/24/98	333	RTA00000427F.k.21.1	M00004090D:F12	62880
3796	3/24/98	130	RTA00000427F.h.12.1	M00004092C:D08	36894
3797	3/24/98	243	RTA00000424F.c.15.3	M00001476D:F12	73533
3798	3/24/98	227	RTA00000427F.i.11.1	M00004097C:H08	26635
3799	3/24/98	456	RTA00000427F.a.10.1	M00004038B:D01	65370
3800	3/24/98	7	RTA00000523F.o.20.1	M00005177B:H02	0
3801	3/24/98	291	RTA00000523F.o.23.1	M00005177C:G04	0
3802	3/24/98	119	RTA00000523F.p.06.1	M00005177D:F09	0
3803	3/24/98	178	RTA00000428F.a.12.1	M00005179B:H02	0
3804	3/24/98	463	RTA00000523F.p.16.1	M00005179D:B03	0
3805	3/24/98	390	RTA00000524F.a.11.1	M00005210D:C09	0
3806	3/24/98	468	RTA00000524F.a.18.1	M00005211A:E09	0
3807	3/24/98	114	RTA00000524F.a.23.1	M00005211C:E09	0
3808	3/24/98	29	RTA00000524F.b.03.1	M00005212A:D10	0
3809	3/24/98	36	RTA00000428F.a.16.1	M00005212D:F08	0
3810	3/24/98	417	RTA00000524F.b.10.1	M00005213C:A01	0
3811	3/24/98	182	RTA00000524F.b.17.1	M00005214B:A06	0
3812	3/24/98	348	RTA00000427F.i.09.1	M00004097C:E03	65916
3813	3/24/98	384	RTA00000527F.p.03.1	M00004029B:A06	5940
3814	3/24/98	84	RTA00000527F.k.18.1	M00003982B:C10	11332
3815	3/24/98	48	RTA00000527F.k.21.1	M00003982B:H10	36051
3816	3/24/98	271	RTA00000527F.1.05.1	M00003983A:D02	13016
3817	3/24/98	246	RTA00000426F.m.21.1	M00003983A:F06	64915
3818	3/24/98	16	RTA00000426F.m.22.1	M00003983A:G02	30002
3819	3/24/98	367	RTA00000527F.I.19.1	M00003983D:E08	36856
3820	3/24/98	477	RTA00000527F.1.21.1	M00003983D:H02	36439
3821	3/24/98	126	RTA00000527F.m.05.1	M00003985A:C01	17240
3822	3/24/98	89	RTA00000527F.n.02.1	M00003986C:G11	24190
3823	3/24/98	263	RTA00000527F.n.07.1	M00003986D:H12	15939
3824	3/24/98	49	RTA00000527F.n.22.1	M00004027A:A08	24175
3825	3/24/98	449	RTA00000426F.m.04.1	M00004028A:B10	36865
3826	3/24/98	336	RTA00000426F.n.17.1	M00004039D:B10	66572
3827	3/24/98	27	RTA00000527F.p.02.1	M00004029B:A01	36844
3828	3/24/98	297	RTA00000525F.c.11.1	M00004039C:E02	37895
3829	3/24/98	17	RTA00000527F.p.06.1	M00004029B:G10	1292
3830	3/24/98	310	RTA00000527F.p.08.1	M00004029C:F02	36013
3831	3/24/98	478	RTA00000527F.p.09.1	M00004029C:F05	7694
3832	3/24/98	253	RTA00000426F.m.08.1	M00004030B:A12	63781
3833	3/24/98	414	RTA00000426F.m.12.1	M00004030B:D08	63740
3834	3/24/98	345	RTA00000426F.n.23.1	M00004030C:A08	18176
3835	3/24/98	98	RTA00000527F.p.16.1	M00004030C:C02	23798
3836	3/24/98	115	RTA00000525F.b.05.1	M00004034C:F05	21116
3837	3/24/98	444	RTA00000525F.b.09.1	M00004035B:F05	23472
3838	3/24/98	158	RTA00000427F.a.06.1	M00004036A:A11	66550
3839	3/24/98	376	RTA00000525F.b.21.1	M00004037C:D04	9486
3840	3/24/98	293	RTA00000525F.c.02.1	M00004038A:E05	14618
3841	3/24/98	138	RTA00000527F.c.22.1	M00003822B:G12	37496

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
3842	3/24/98	30	RTA00000426F.m.07.1	M00004028A:G03	63504
3843	3/24/98	322	RTA00000523F.i.17.1	M00003856B:A12	65779
3844	3/24/98	311	RTA00000428F.b.02.1	M00005214D:D10	0
3845	3/24/98	233	RTA00000426F.f.13.1	M00003851A:A06	65384
3846	3/24/98	274	RTA00000523F.h.06.1	M00003851B:D03	28745
3847	3/24/98	407	RTA00000523F.h.08.1	M00003851B:E01	62893
3848	3/24/98	82	RTA00000523F.h.15.1	M00003851C:F09	65137
3849	3/24/98	316	RTA00000523F.h.16.1	M00003851D:H11	66031
3850	3/24/98	232	RTA00000527F.i.13.2	M00003852B:G04	2924
3851	3/24/98	451	RTA00000527F.i.15.2	M00003852C:F07	14235
3852	3/24/98	249	RTA00000523F.h.21.1	M00003853B:C10	41440
3853	3/24/98	72	RTA00000426F.f.19.1	M00003854C:C09	66701
3854	3/24/98	60	RTA00000523F.i.06.1	M00003855A:A01	66341
3855	3/24/98	91	RTA00000527F.i.21.2	M00003855A:F01	37490
3856	3/24/98	137	RTA00000527F.h.17.1	M00003848D:G02	37799
3857	3/24/98	433	RTA00000527F.j.04.2	M00003856A:G04	11809
3858	3/24/98	157	RTA00000523F.g.10.1	M00003848B:E07	40694
3859	3/24/98	75	RTA00000523F.i.22.1	M00003857A:E12	64688
3860	3/24/98	481	RTA00000523F.j.02.1	M00003857A:H10	62853
3861	3/24/98	377	RTA00000527F.j.12.2	M00003857C:E05	37503
3862	3/24/98	286	RTA000003277 g.12.2	M00003858B:G02	63672
3863	3/24/98	71	RTA00000527F.j.20.2	M00003860D:E06	37603
3 8 64	3/24/98	205	RTA00000426F.h.12.1	M00003905C:F12	78093
3865	3/24/98	40	RTA00000426F.h.23.1	M00003911A:D12	75964
3866	3/24/98	369	RTA00000524F.c.08.1	M00005217C:C01	0
3867	3/24/98	234	RTA00000524F.c.16.1	M00005218D:G10	0
3868	3/24/98	8	RTA00000428F.b.06.1	M00005228A:A09	0
3869	3/24/98	193	RTA00000428F.b.12.1	M00005231C:B07	0
3870	3/24/98	419	RTA00000428F.b.22.1	M00005231D:B09	0
3871	2/24/98	486	RTA00000346F.I.13.1	M00003980B:C11	7542
3872	3/24/98	421	RTA00000523F.i.10.1	M00003855B:B09	64876
3873	3/24/98	10	RTA00000527F.f.12.1	M00003829D:D12	5945
3874	3/24/98	145	RTA00000426F.m.24.1	M00003981A:A07	63943
3875	3/24/98	39	RTA00000527F.c.23.1	M00003822C:A07	37742
3876	3/24/98	35	RTA00000426F.f.11.1	M00003823C:B01	63102
3877	3/24/98	385	RTA00000426F.f.12.1	M00003823C:C04	19096
3878	3/24/98	2	RTA00000523F.d.19.1	M00003824A:A06	26489
3879	3/24/98	225	RTA00000527F.d.09.1	M00003824A:G11	10848
3880	3/24/98	359	RTA00000523F.d.21.1	M00003824B:C09	33424
3881	3/24/98	330	RTA00000523F.d.23.1	M00003824C:A10	63633
3882	3/24/98	254	RTA00000523F.d.24.1	M00003824D:D08	64799
3883	3/24/98	74	RTA00000527F.d.19.1	M00003825B:F10	486
3884	3/24/98	67	RTA00000527F.e.03.1	M00003825D:F01	25560
3885	3/24/98	352	RTA00000527F.e.13.1	M00003826C:F05	37588
3886	3/24/98	185	RTA00000527F.h.21.1	M00003850C:G09	37630
3887	3/24/98	338	RTA00000523F.e.15.1	M00003829C:E08	7919
3888	3/24/98	284	RTA00000524F.b.19.1	M00005216B:D02	0
3889	3/24/98	242	RTA00000523F.e.20.1	M00003829D:F03	65164
3890	3/24/98	301	RTA00000527F.f.18.1	M00003830D:B11	37577
3891	3/24/98	259	RTA00000528F.m.04.1	M00003830D:H11	10815

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
2002	Appln	Appln	DEL 00000500E 006 1		
3892	3/24/98	160	RTA00000523F.f.06.1	M00003833D:H08	62871
3893	3/24/98	166	RTA00000523F.f.07.1	M00003833D:H10	62799
3894	3/24/98	196	RTA00000523F.f.12.1	M00003840A:C10	63751
3895	3/24/98	447	RTA00000523F.f.19.1	M00003840B:F05	34169
3896	3/24/98	113	RTA00000527F.g.07.1	M00003840C:C02	37488
3897	3/24/98	45	RTA00000528F.m.12.1	M00003842D:F08	5768
3898	3/24/98	415	RTA00000527F.g.12.1	M00003845C:D04	37746
3899	3/24/98	1	RTA00000527F.g.13.1	M00003845D:A04	36035
3900	3/24/98	450	RTA00000527F.g.21.1	M00003846B:C05	36028
3901	3/24/98	144	RTA00000527F.g.23.1	M00003846C:F08	37538
3902	3/24/98	422	RTA00000527F.f.03.1	M00003829A:B08	17788
3903	3/24/98	176	RTA00000522F.g.15.1	M00001595B:G07	76536
3904	3/24/98	264	RTA00000425F.e.09.1	M00001608C:G04	75550
3905	3/24/98	32	RTA00000424F.n.14.1	M00001584D:C11	73008
3906	3/24/98	459	RTA00000425F.c.03.1	M00001585D:B12	74643
3907	3/24/98	124	RTA00000424F.m.12.1	M00001586C:H07	77675
3908	3/24/98	314	RTA00000522F.e.09.1	M00001589D:A01	32599
3909	3/24/98	262	RTA00000424F.k.03.1	M00001590D:B04	21289
3910	3/24/98	106	RTA00000424F.j.14.1	M00001592B:B02	74311
3911	3/24/98	424	RTA00000424F.k.10.1	M00001592D:H02	73232
3912	3/24/98	168	RTA00000424F.j.12.1	M00001594C:E05	73827
3913	3/24/98	420	RTA00000424F.j.13.1	M00001594C:H03	74485
3914	3/24/98	210	RTA00000522F.g.06.1	M00001594D:G11	78221
3915	3/24/98	439	RTA00000522F.g.10.1	M00001595A:C07	74294
3916	3/24/98	300	RTA00000424F.m.08.1	M00001584A:A07	19402
3917	3/24/98	355	RTA00000522F.g.12.1	M00001595A:E07	78783
3918	3/24/98	411	RTA00000424F.n.12.1	M00001582C:G02	41589
3919	3/24/98	238	RTA00000522F.g.17.1	M00001595B:G10	76486
3920	3/24/98	472	RTA00000522F.g.18.1	M00001595B:H11	73226
3921	3/24/98	102	RTA00000522F.g.19.1	M00001595C:A01	78119
3922	3/24/98	280	RTA00000522F.g.20.1	M00001595C:A05	77688
3923	3/24/98	191	RTA00000522F.g.22.1	M00001595C:B12	77504
3924	3/24/98	163	RTA00000522F.h.01.1	M00001595C:E05	75010
3925	3/24/98	438	RTA00000522F.h.02.1	M00001595C:E09	74947
3926	3/24/98	257	RTA00000522F.h.07.1	M00001595D:C11	75149
3927	3/24/98	389	RTA00000424F.i.15.1	M00001596A:A02	78043
3928	3/24/98	167	RTA00000424F.i.20.1	M00001596A:D01	44010
3929	3/24/98	374	RTA00000528F.f.10.1	M00001596C:G05	3600
3930	3/24/98	215	RTA00000425F.f.02.1	M00001607A:A01	76982
3931	3/24/98	22	RTA00000527F.k.15.1	M00003982A:G03	22688
3932	3/24/98	378	RTA00000522F.g.11.1	M00001595A:D12	75432
3933	3/24/98	63	RTA00000522F.b.01.1	M00001570C:B02	75691
3934	3/24/98	430	RTA00000424F.g.08.1	M00001482C:F09	74928
3935	3/24/98	340	RTA00000424F.h.06.1	M00001485C:D07	77552
3936	3/24/98	161	RTA00000424F.h.10.1	M00001485C:G06	72925
3937	3/24/98	368	RTA00000424F.i.11.1	M00001485D:A05	41569
3938	3/24/98	455	RTA00000424F.g.24.1	M00001487C:A11	79156
3939	3/24/98	211	RTA00000424F.h.03.1	M00001487C:G09	74447
3940	3/24/98	174	RTA00000424F.b.21.4	M00001530A:B02	24686
3941	3/24/98	383	RTA00000424F.b.23.4	M00001530A:H05	77322

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
2042	Appln 3/24/98	Appln 179	RTA00000424F.d.10.3	M00001530D:A11	73110
3942 3943	3/24/98 3/24/98	393	RTA00000424F.b.15.4	M00001530B:R11	74958
	3/24/98	343 347	RTA000004241.0.15.4 RTA00000522F.a.05.1	M00001557B:D10	32611
3944		303	RTA00000522F.a.06.1	M00001567A:C04	73662
3945	3/24/98	303 229	RTA00000322F.a.00.1 RTA00000424F.n.13.1	M00001584D:B06	74942
3946	3/24/98		RTA00000424F.ii.13.1 RTA00000522F.a.20.1	M00001567C:E07	74070
3947	3/24/98	392		M00001507C.E07	75921
3948	3/24/98	226	RTA00000425F.e.15.1	M00001008D:111 M00001570D:E05	78634
3949	3/24/98	285	RTA00000522F.b.07.1	M00001570D:E03 M00001570D:E07	2395
3950	3/24/98	465	RTA00000528F.d.04.1	M00001570D:E07 M00001573B:A06	3460
3951	3/24/98	404	RTA00000522F.b.18.1	M00001573B:A00 M00001573B:H12	75181
3952	3/24/98	9	RTA00000522F.b.22.1	M00001575B:H12	43214
3953	3/24/98	109	RTA00000424F.a.01.4	M00001575A:D05	43214
3953	3/24/98	125	RTA00000424F.a.01.1		43214
3954	3/24/98	125	RTA00000424F.a.01.1	M00001575A:D05	43214
3954	3/24/98	109	RTA00000424F.a.01.4	M00001575A:D05	
3955	3/24/98	294	RTA00000424F.a.05.1	M00001575B:C01	77976
3955	3/24/98	292	RTA00000424F.a.05.4	M00001575B:C01	77976
3956	3/24/98	292	RTA00000424F.a.05.4	M00001575B:C01	77976
3956	3/24/98	294	RTA00000424F.a.05.1	M00001575B:C01	77976
3957	3/24/98	434	RTA00000522F.c.11.1	M00001576C:H02	31064
3958	3/24/98	299	RTA00000522F.c.14.1	M00001577A:A03	75449
3959	3/24/98	110	RTA00000522F.d.08.1	M00001578B:A06	74284
3960	3/24/98	306	RTA00000522F.d.23.1	M00001579D:F02	73868
3961	3/24/98	350	RTA00000424F.n.11.1	M00001582C:C04	73874
3962	3/24/98	366	RTA00000522F.a.17.1	M00001567C:B08	79032
3963	3/24/98	239	RTA00000523F.j.17.1	M00003966B:A04	63610
3964	3/24/98	405	RTA00000425F.e.07.1	M00001608C:D02	75992
3965	3/24/98	231	RTA00000426F.e.17.1	M00003810C:B06	64089
3966	3/24/98	104	RTA00000527F.b.18.1	M00003810D:H09	37469
3967	3/24/98	312	RTA00000426F.f.17.1	M00003811C:C02	66334
3968	3/24/98	266	RTA00000426F.f.16.1	M00003813B:F02	65613
3969	3/24/98	183	RTA00000527F.c.04.1	M00003813C:H08	23090
3970	3/24/98	435	RTA00000523F.c.13.1	M00003813D:B12	40668
3971	3/24/98	255	RTA00000523F.c.14.1	M00003813D:C02	66015
3972	3/24/98	131	RTA00000523F.c.15.1	M00003813D:G06	36935
3973	3/24/98	270	RTA00000426F.g.16.1	M00003814B:C01	41446
3974	3/24/98	95	RTA00000523F.c.18.1	M00003817C:A10	66179
3975	3/24/98	329	RTA00000527F.c.09.1	M00003817C:G06	64859
3976	3/24/98	65	RTA00000523F.c.01.1	M00003810A:A02	65710
3977	3/24/98	398	RTA00000527F.c.16.1	M00003821A:H09	22908
3978	3/24/98	96	RTA00000523F.b.13.1	M00003809B:A03	66330
3979	3/24/98	313	RTA00000523F.j.21.1	M00003966C:A12	36925
3980	3/24/98	86	RTA00000523F.k.01.1	M00003966C:F03	41437
3981	3/24/98	26	RTA00000427F.b.23.1	M00003973D:F08	64297
3982	3/24/98	277	RTA00000427F.e.08.1	M00003974D:E01	47387
3983	3/24/98	397	RTA00000427F.e.10.1	M00003974D:H07	64599
3984	3/24/98	31	RTA00000427F.c.10.1	M00003976B:E06	65478
3985	3/24/98	151	RTA00000427F.c.12.1	M00003976B:H07	66995
3986	3/24/98	57	RTA00000427F.c.20.1	M00003978A:E01	26527
3987	3/24/98	213	RTA00000427F.c.22.1	M00003978A:E09	63990

Appln Appln Appln 3988 374/98 289 RTA00000427F.d.10.1 M00003978C:A12 40685 3989 3724/98 28 RTA00000427F.d.08.1 M00003980C:E12 63967 3990 3724/98 335 RTA00000427F.d.09.1 M00003980C:E12 63967 3724/98 335 RTA00000427F.d.10.9.1 M00003980C:E12 63967 3724/98 343 RTA00000527F.c.11.1 M00001635B:B02 75305 3992 3724/98 343 RTA00000527F.c.11.1 M00001650C-Q4 40841 3994 3724/98 155 RTA00000424F.I.21.1 M00001650C-Q4 40841 3994 3724/98 312 RTA00000424F.I.21.1 M00001609C:G05 79017 3995 3724/98 214 RTA00000424F.I.21.1 M00001609C:G05 79017 3996 3724/98 214 RTA00000424F.I.21.1 M00001653D:H07 74071 3998 3724/98 428 RTA00000522F.I.08.1 M000016540C:B07 73661 4000 3724/98 275 RTA00000522F.I.08.1 M00001654A:E08 78781 3999 3724/98 223 RTA00000522F.I.02.1 M00001654C:G07 75834 4002 3724/98 391 RTA00000522F.m.02.1 M00001654C:G07 75834 4002 3724/98 346 RTA00000522F.m.02.1 M00001655C:C07 41544 4004 3724/98 332 RTA00000522F.m.03.1 M00001655C:C07 41544 4006 3724/98 332 RTA00000522F.m.03.1 M00001655C:C07 41544 4006 3724/98 51 RTA00000522F.m.03.1 M00001655C:C07 41544 4006 3724/98 58 RTA00000522F.m.05.1 M00001655D:E08 74959 4006 3724/98 58 RTA00000522F.m.05.1 M00001655D:E08 74959 4006 3724/98 58 RTA00000522F.m.05.1 M00001655D:E08 74959 4008 3724/98 58 RTA00000522F.m.05.1 M00001655D:D111 3260 4007 3724/98 58 RTA00000522F.m.05.1 M00001650C:C04 79275 4008 3724/98 58 RTA0000052F.m.05.1 M00001655D:D111 3724/98 50 RTA0000052F.m.05.1 M00001650C:C04 79275 4008 3724/98 58 RTA0000052F.m.10.1 M00001650C:C04 79275 4008 3724/98 58 RTA0000052F.m.1	SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
3989 3/24/98 335	***			D.T. 1 00000 10 T. 1 1 1 1 1 1		
3990 3/24/98 267						
3991 3/24/98 343 RTA00000527F.c.11.1 M00001635B:B02 75305 37992 3724/98 343 RTA00000527F.c.11.1 M00001656B:B02 37484 3994 3724/98 351 RTA00000424F.I19.1 M00001609C:A12 75454 3995 3724/98 321 RTA00000424F.I19.1 M00001609C:A12 75454 3995 3724/98 321 RTA00000424F.in.04.1 M00001609C:A12 75665 3997 3724/98 321 RTA00000424F.k.12.1 M00001609C:A12 77666 3997 3724/98 446 RTA00000425F.f.20.1 M00001650E:B07 776766 3997 3724/98 428 RTA00000522F.in.8.1 M00001654B:A01 74671 4000 3724/98 295 RTA00000522F.in.8.1 M00001654B:A01 74691 4000 3724/98 223 RTA00000522F.in.02.1 M00001654C:D10 75801 4001 3724/98 391 RTA00000522F.in.02.1 M00001654C:D10 75801 4002 3724/98 391 RTA00000522F.in.03.1 M00001654C:C07 76834 4004 3724/98 391 RTA00000522F.in.02.1 M00001655C:C07 41544 4004 3724/98 394 RTA00000522F.in.02.1 M00001655C:C07 41544 4004 3724/98 394 RTA00000522F.in.02.1 M00001655D:D68 74959 4005 3724/98 332 RTA00000522F.in.05.1 M00001655D:H11 73260 4006 3724/98 332 RTA00000522F.in.05.1 M00001655D:H11 73260 4006 3724/98 58 RTA00000522F.in.05.1 M00001655D:H11 73260 4008 3724/98 58 RTA00000522F.in.6.1 M00001657D:A10 26769 4011 3724/98 20 RTA00000522F.in.6.1 M00001657D:A10 26769 4011 3724/98 38 RTA0000052F.in.6.1 M00001657D:A10 26769 4011 3724/98 341 RTA0000052F.in.6.1 M00001659D:A09 26860 4012 3724/98 341 RTA0000052F.in.6.1 M00001659D:A09 26860 4014 3724/98						
3992 3/24/98 251						
3993 3/24/98 251						
3994 3/24/98 155						
3995 3/24/98 321 RTA00000424F.m.04.1 M00001609C:G05 79017 3996 3/24/98 214 RTA00000424F.k.12.1 M00001610C:B07 77667 3997 3/24/98 428 RTA00000522F.L08.1 M00001653D:H07 74071 3998 3/24/98 295 RTA00000522F.L08.1 M00001654A:E08 78781 3999 3/24/98 225 RTA00000522F.I.2.1 M00001654C:G07 76834 4001 3/24/98 223 RTA00000522F.m.02.1 M00001654C:G07 76834 4002 3/24/98 391 RTA00000522F.m.03.1 M00001655C:C07 76834 4004 3/24/98 391 RTA00000522F.m.02.1 M00001655C:C09 7994 4005 3/24/98 34 RTA00000522F.m.05.1 M00001655D:E08 74959 4006 3/24/98 31 RTA00000522F.m.05.1 M00001655D:E08 74959 4007 3/24/98 31 RTA00000522F.m.05.1 M00001657C:C11 73691 4007 3/24/98 172						
3996 3/24/98 446 RTA00000424F.k.12.1 M00001610C:B07 77666 3997 3/24/98 446 RTA00000425F.f.20.1 M00001653D:H07 74071 3998 3/24/98 295 RTA00000522F.l.08.1 M00001654C:B08 78781 3999 3/24/98 295 RTA00000522F.l.15.1 M00001654C:D10 75801 4001 3/24/98 223 RTA00000522F.m.02.1 M00001654C:G09 79194 4002 3/24/98 391 RTA00000522F.m.02.1 M00001654C:G09 79194 4003 3/24/98 391 RTA00000522F.m.02.1 M00001654C:G09 79194 4003 3/24/98 31 RTA00000522F.m.02.1 M00001655C:C07 41544 4004 3/24/98 51 RTA00000522F.m.02.1 M00001655D:E08 74959 4005 3/24/98 32 RTA00000522F.m.05.1 M00001655D:E08 74959 4006 3/24/98 32 RTA00000522F.n.05.1 M00001655D:H11 73260 4006 3/24/98 172 RTA00000522F.n.05.1 M00001655D:H11 73260 4006 3/24/98 58 RTA00000522F.n.06.1 M00003810B:B1 212469 4009 3/24/98 56 RTA00000522F.n.16.1 M00001657D:A10 26769 4010 3/24/98 20 RTA00000522F.n.16.1 M00001657D:A10 26769 4011 3/24/98 20 RTA00000522F.n.06.1 M00001657D:A10 26769 4013 3/24/98 413 RTA00000522F.n.06.1 M00001668B:D05 2478 4013 3/24/98 413 RTA00000522F.n.06.1 M00001669D:D05 2478 4013 3/24/98 38 RTA00000522F.n.06.1 M00001669D:D05 2478 4013 3/24/98 38 RTA00000522F.n.06.1 M00001669D:D05 2478 4014 3/24/98 413 RTA00000522F.n.06.1 M00001669B:C12 1070 4016 3/24/98 31 RTA00000522F.n.06.1 M00001669B:C12 1070 4016 3/24/98 31 RTA00000522F.n.01 M00001669B:C12 1070 4016 3/24/98 31 RTA00000522F.n.01 M00001669B:C12 1070 4016 3/24/98 31 RTA00000522F.n.01 M00001669B:C12 1070 4016 3/24/98 31 RTA00000527F.a.13.1 M00001669B:C12 1070 4016 3/24/98 31 RTA00000527F.n.13.1 M00001669B:C12 1070 4018 3/24/98 31 RTA00000527F.n.13.1 M00001669B:C10 37740 4020 3/24/98 356 RTA00000527F.n.13.1 M000016708:CP09 1981 4019 3/24/98 360 RTA00000527F.n.13.1 M000016708:CP09 1981 4020 3/24/98 31 RTA0000049F.c.00.1 M000016708:CP09 1981 4021 3/24/98 31 RTA0000049F.c.00.1 M000016708:CP09 1981 4022 3/24/98 1181 RTA0000049F.c.00.1 M00003802B:D10 21421 4024 2/24/98 1181 RTA0000049F.c.01 M0000382D:D00 66065 4022 3/24/98 528 RTA0000049F.c.16.1 M0000382D:D00 66495 4033 2/24/98 81 RTA00000419F.c.16.1 M0000383D:C003 65963						
3997 3/24/98 446 RTA00000425F.f.20.1 M00001653D:H07 74071 3998 3/24/98 428 RTA00000522F.l.08.1 M00001654A:E08 78781 3999 3/24/98 295 RTA00000522F.l.15.1 M00001654C:G07 74691 4000 3/24/98 275 RTA00000522F.m.02.1 M00001654C:G07 76834 4002 3/24/98 391 RTA00000522F.m.02.1 M00001654C:G07 76834 4003 3/24/98 391 RTA00000522F.m.03.1 M00001655C:C07 41544 4004 3/24/98 31 RTA00000522F.m.02.1 M00001655C:C07 41544 4004 3/24/98 51 RTA00000522F.m.02.1 M00001655D:E08 74959 4005 3/24/98 94 RTA00000522F.m.05.1 M00001655D:E08 74959 4006 3/24/98 332 RTA00000522F.m.05.1 M00001655D:B11 36913 4007 3/24/98 172 RTA00000522F.m.05.1 M00001656C:C04 79275 4008 3/24/98 58 RTA00000522F.m.14.1 M00001656C:C04 79275 4008 3/24/98 58 RTA00000522F.m.14.1 M00001657D:A10 26769 4011 3/24/98 20 RTA00000522F.n.16.1 M00001657D:A10 26769 4011 3/24/98 30 RTA00000522F.n.16.1 M00001657D:A00 26860 4012 3/24/98 38 RTA00000522F.n.16.1 M00001657D:A00 26860 4012 3/24/98 412 RTA00000425F.i.10.1 M00001669D:A09 26860 4012 3/24/98 413 RTA00000425F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000425F.i.11.1 M00001664B:E08 78736 4016 3/24/98 432 RTA00000522F.n.06.1 M00001669C:D0 2478 4018 3/24/98 432 RTA00000522F.n.06.1 M00001669B:C10 1070 4018 3/24/98 31 RTA00000522F.n.00.1 M00001669B:C10 1070 4018 3/24/98 31 RTA00000522F.n.00.1 M00001669B:C10 1070 4018 3/24/98 31 RTA00000522F.n.00.1 M00001669B:C10 1070 4018 3/24/98 31 RTA00000522F.n.0.1 M00001669B:C10 1700 4018 3/24/98 31 RTA00000522F.n.0.1 M00001669B:C10 1700 4019 3/24/98 31 RTA00000522F.n.0.1 M00001669B:C10 1700 4018 3/24/98 31 RTA0000052F.n.0.1 M00001669C:D00 74853 4017 3/24/98 31 RTA00000419F.c.10.1 M00001670A:F09 75204 4018 3/24/98 31 RTA00000409F.n.03.1 M0000382D:D06 37740 4021 3/24/98 1181 RTA00000409F.n.03.1 M0000382D:D06 37540 4022 3/24/98 1181 RTA00000419F.c.02.1 M0000382D:D04 45815 4024 2/24/98 1181 RTA00000419F.c.06.1 M0000382D:D04 45815 4025 2/24/98 1181 RTA00000419F.c.16.1 M0000382D:D04 45815 4033 2/24/98 81 RTA00000419F.c.16.1 M0000383D:C05 62963						
3998 3/24/98 428 RTA00000522F.I.08.1 M00001654A:E08 78781 3999 3/24/98 295 RTA00000522F.I.15.1 M00001654B:A01 74691 4000 3/24/98 223 RTA00000522F.m.02.1 M00001654C:G07 76834 4001 3/24/98 391 RTA00000522F.m.03.1 M00001654C:G09 79194 4003 3/24/98 391 RTA00000522F.m.03.1 M00001655C:C07 41544 4004 3/24/98 51 RTA00000522F.m.02.1 M00001655D:E08 74959 4005 3/24/98 31 RTA00000522F.m.05.1 M00001655D:E08 74959 4006 3/24/98 32 RTA00000522F.m.05.1 M00001655D:H11 73260 4006 3/24/98 332 RTA00000522F.m.05.1 M00001655D:H11 73260 4006 3/24/98 332 RTA00000522F.m.05.1 M00001655D:H11 73260 4007 3/24/98 172 RTA00000522F.m.05.1 M00001655D:H11 73260 4008 3/24/98 58 RTA00000522F.m.14.1 M00001657C:C17 73410 4010 3/24/98 56 RTA00000522F.m.16.1 M00001657C:C11 73410 4011 3/24/98 20 RTA00000522F.m.16.1 M00001657D:A10 26769 4011 3/24/98 38 RTA00000522F.m.16.1 M00001657D:A10 26769 4011 3/24/98 38 RTA00000522F.m.16.1 M00001657D:A10 26769 4011 3/24/98 412 RTA00000522F.m.10.1 M00001669D:A09 26860 4012 3/24/98 412 RTA00000522F.m.10.1 M00001669B:C12 1070 4016 3/24/98 420 RTA00000522F.m.00.1 M00001669B:C12 1070 4016 3/24/98 31 RTA00000522F.m.01 M00001669B:C12 1070 4018 3/24/98 331 RTA00000522F.n.01 M00001669B:C12 1070 4019 3/24/98 202 RTA00000522F.n.01 M00001669B:C12 1070 4018 3/24/98 331 RTA00000522F.n.01 M00001669B:C12 1070 4019 3/24/98 336 RTA00000522F.n.01 M00001669B:C12 1070 4019 3/24/98 331 RTA00000522F.n.01 M00001670A:F09 75204 4018 3/24/98 331 RTA00000522F.n.01 M00001670A:F09 75204 4018 3/24/98 31 RTA00000522F.n.01 M00001670B:D66 37740 4021 3/24/98 118 RTA0000052F.n.01 M00001670B:D66 37740 4021 3/24/98 12 RTA0000052F.n.01 M00001670B:D66 37740 4021 3/24/98 356 RTA0000052F.n.01 M00001670B:D66 37740 4021 3/24/98 14 RTA0000052F.n.03.1 M0000380B:F03 16998 4025 2/24/98 118 RTA0000049F.c.02.1 M0000382D:D10 21421 4026 2/24/98 118 RTA00000409F.n.03.1 M0000382D:D00 45815 4024 2/24/98 115 RTA00000419F.c.05.1 M0000382D:D10 65254 4033 2/24/98 528 RTA00000419F.c.16.1 M0000382D:D04 65254 4035 2/24/98 81 RTA00000419F.c.16.1 M0000383D:C036 65963						
3999 3/24/98 295 RTA00000522F.I.15.1 M00001654B:A01 74691 4000 3/24/98 275 RTA00000522F.m.02.1 M00001654C:B10 75801 4001 3/24/98 391 RTA00000522F.m.03.1 M00001654C:G09 79194 4003 3/24/98 346 RTA00000522F.m.03.1 M00001655D:C07 41544 4004 3/24/98 51 RTA00000522F.m.05.1 M00001655D:E08 74959 4005 3/24/98 94 RTA00000522F.m.05.1 M00001655D:H11 73260 4006 3/24/98 32 RTA00000522F.m.05.1 M00001656C:C04 79275 4008 3/24/98 172 RTA00000522F.n.16.1 M00001656C:C04 79275 4009 3/24/98 28 RTA00000522F.n.14.1 M00001657D:A10 26769 4010 3/24/98 20 RTA00000522F.n.16.1 M00001657D:A10 26769 4011 3/24/98 38 RTA00000522F.o.06.1 M00001669D:A0 72478 4013 3/24/98 38					M00001653D:H07	
4000 3/24/98 275 RTA00000522F.I.22.1 M00001654C:D10 75801 4001 3/24/98 223 RTA00000522F.m.02.1 M00001654C:G09 79194 4002 3/24/98 391 RTA00000522F.m.03.1 M00001654C:G09 79194 4003 3/24/98 346 RTA00000522F.m.02.1 M00001655C:C07 41544 4004 3/24/98 51 RTA00000522F.n.02.1 M00001655D:E08 74959 4006 3/24/98 94 RTA00000522F.n.03.1 M00001655D:D111 73260 4006 3/24/98 172 RTA00000522F.n.03.1 M00001655D:D111 73260 4008 3/24/98 18 RTA00000522F.n.06.1 M00001657C:C11 73410 4009 3/24/98 20 RTA00000522F.n.14.1 M00001657C:C11 73410 4011 3/24/98 20 RTA00000522F.o.06.1 M00001659D:A09 26860 4012 3/24/98 38 RTA0000052EF.i.22.1 M00001664B:E08 78736 4013 3/24/98 412						
4001 3/24/98 223 RTA00000522F.m.02.1 M00001654C:G07 76834 4002 3/24/98 391 RTA00000522F.m.03.1 M00001655C:C07 41544 4003 3/24/98 51 RTA00000522F.m.02.1 M00001655D:E08 74959 4005 3/24/98 94 RTA00000522F.n.05.1 M00001655D:E08 74959 4006 3/24/98 332 RTA00000522F.n.05.1 M00001655D:E08 74959 4006 3/24/98 312 RTA000000522F.n.05.1 M0000185B:B11 36913 4007 3/24/98 172 RTA000000522F.n.06.1 M00001657C:C01 79275 4008 3/24/98 58 RTA00000522F.n.16.1 M00001657D:A10 26769 4010 3/24/98 26 RTA00000522F.o.06.1 M00001657D:A10 26769 4011 3/24/98 38 RTA00000522F.o.06.1 M00001657D:A10 26769 4012 3/24/98 38 RTA00000425F.i.10.1 M00001669D:D05 2478 4013 3/24/98 412					M00001654B:A01	74691
4002 3/24/98 391 RTA00000522F.m.03.1 M00001654C:G09 79194 4003 3/24/98 346 RTA00000522F.m.19.1 M00001655C:C07 41544 4004 3/24/98 51 RTA00000522F.n.05.1 M00001655D:H11 73260 4006 3/24/98 94 RTA00000523F.c.03.1 M00001655D:H11 73260 4006 3/24/98 172 RTA00000523F.c.03.1 M00001656C:C04 79275 4008 3/24/98 172 RTA00000527F.k.06.1 M00003981BB12 12469 4009 3/24/98 240 RTA00000522F.n.16.1 M00001657C:C11 73410 4010 3/24/98 240 RTA00000522F.n.16.1 M00001657D:A10 26769 4011 3/24/98 20 RTA00000522F.o.06.1 M00001661D:D05 2478 4013 3/24/98 413 RTA00000522F.o.06.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000528F.i.10.1 M00001669B:C12 1070 4016 3/24/98 412				RTA00000522F.1.22.1	M00001654C:D10	75801
4003 3/24/98 346 RTA00000522F.m.19.1 M00001655C:C07 41544 4004 3/24/98 51 RTA00000522F.n.02.1 M00001655D:E08 74959 4005 3/24/98 332 RTA00000522F.n.03.1 M00003810B:B11 36913 4007 3/24/98 332 RTA0000052F.b.06.1 M00003810B:B11 36913 4008 3/24/98 58 RTA0000052F.b.06.1 M0000381B:B12 12469 4009 3/24/98 58 RTA0000052F.n.14.1 M00001657C:C11 73410 4010 3/24/98 20 RTA0000052F.n.16.1 M00001657D:A10 26769 4011 3/24/98 38 RTA0000052F.n.06.1 M00001659D:A09 26860 4012 3/24/98 38 RTA0000052F.n.06.1 M00001661D:D05 2478 4013 3/24/98 38 RTA0000052F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA0000052F.i.10.1 M00001669B:C12 1070 4016 3/24/98 422 RTA0	4001	3/24/98		RTA00000522F.m.02.1	M00001654C:G07	76834
4004 3/24/98 51 RTA00000522F.n.02.1 M00001655D:E08 74959 4005 3/24/98 94 RTA00000522F.n.05.1 M00001655D:H11 73260 4006 3/24/98 332 RTA00000523F.c.03.1 M00003810B:B11 36913 4007 3/24/98 172 RTA00000425F.f.11.1 M00001656C:C04 79275 4008 3/24/98 58 RTA00000522F.n.16.1 M00003981B:B12 12469 4009 3/24/98 240 RTA00000522F.n.16.1 M000016570:A10 26769 4011 3/24/98 20 RTA00000522F.n.16.1 M00001659D:A09 26860 4012 3/24/98 38 RTA00000525F.i.10.1 M00001669D:A09 26860 4013 3/24/98 413 RTA00000525F.i.10.1 M00001661D:D05 2478 4014 3/24/98 412 RTA000000525F.i.10.1 M00001669B:C01 1070 4016 3/24/98 432 RTA00000525F.j.11.1 M00001669C:B09 74853 4017 3/24/98 21	4002	3/24/98	391	RTA00000522F.m.03.1	M00001654C:G09	79194
4005 3/24/98 94 RTA00000522F.n.05.1 M00001655D:H11 73260 4006 3/24/98 332 RTA00000523F.c.03.1 M00003810B:B11 36913 4007 3/24/98 172 RTA00000425F.f.11.1 M00001656C:C04 79275 4008 3/24/98 58 RTA00000522F.n.16.1 M00003981B:B12 12469 4009 3/24/98 240 RTA00000522F.n.16.1 M00001657D:A10 26769 4010 3/24/98 56 RTA00000522F.n.06.1 M00001659D:A09 26860 4011 3/24/98 20 RTA00000528F.i.2.1 M00001659D:A09 26860 4012 3/24/98 38 RTA00000528F.i.10.1 M00001659D:A09 26860 4013 3/24/98 412 RTA00000528F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000528F.j.11.1 M00001669B:C12 1070 4015 3/24/98 432 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 331	4003	3/24/98	346	RTA00000522F.m.19.1	M00001655C:C07	41544
4006 3/24/98 332 RTA00000523F.c.03.1 M00003810B:B11 36913 4007 3/24/98 172 RTA00000425F.f.11.1 M00001656C:C04 79275 4008 3/24/98 58 RTA00000527F.k.06.1 M00003981B:B12 12469 4009 3/24/98 240 RTA00000522F.n.14.1 M00001657C:C11 73410 4010 3/24/98 56 RTA00000522F.n.16.1 M00001657D:A10 26769 4011 3/24/98 20 RTA00000522F.o.06.1 M00001659D:A09 26860 4012 3/24/98 38 RTA00000522F.o.06.1 M00001661D:D05 2478 4013 3/24/98 413 RTA00000425F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000522F.o.20.1 M00001669B:C12 1070 4016 3/24/98 432 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000522F.o.20.1 M00001678C:F09 1981 4019 3/24/98 331	4004	3/24/98	51	RTA00000522F.n.02.1	M00001655D:E08	74959
4007 3/24/98 172 RTA00000425F.f.11.1 M00001656C:C04 79275 4008 3/24/98 58 RTA00000527F.k.06.1 M00003981B:B12 12469 4009 3/24/98 240 RTA00000522F.n.16.1 M00001657C:C11 73410 4010 3/24/98 56 RTA00000522F.n.06.1 M00001657D:A10 26769 4011 3/24/98 20 RTA00000522F.n.06.1 M00001659D:A09 26860 4012 3/24/98 38 RTA00000522F.n.06.1 M00001661D:D05 2478 4013 3/24/98 413 RTA00000528F.i.22.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000528F.i.10.1 M00001669B:C12 1070 4016 3/24/98 432 RTA00000522F.n.20.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000522F.n.20.1 M00001670A:F09 75204 4018 3/24/98 331 RTA00000523F.a.07.1 M00001670A:F09 75204 4019 3/24/98 356	4005	3/24/98	94	RTA00000522F.n.05.1	M00001655D:H11	73260
4008 3/24/98 58 RTA00000527F.k.06.1 M00003981B:B12 12469 4009 3/24/98 240 RTA00000522F.n.14.1 M00001657C:C11 73410 4010 3/24/98 56 RTA00000522F.o.06.1 M00001657D:A10 26769 4011 3/24/98 20 RTA00000522F.o.06.1 M00001657D:A09 26860 4012 3/24/98 38 RTA00000528F.i.22.1 M00001661D:D05 2478 4013 3/24/98 413 RTA00000425F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000528F.i.11.1 M00001669B:C12 1070 4016 3/24/98 432 RTA00000528F.j.11.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000528F.k.10.1 M00001678C:F09 78204 4018 3/24/98 331 RTA00000528F.k.10.1 M00001693A:H06 75804 4020 3/24/98 356 RTA00000528F.k.10.1 M00001866:H12 7740 4021 3/24/98 14	4006	3/24/98	332	RTA00000523F.c.03.1	M00003810B:B11	36913
4009 3/24/98 240 RTA00000522F.n.14.1 M00001657C:C11 73410 4010 3/24/98 56 RTA00000522F.n.16.1 M00001657D:A10 26769 4011 3/24/98 20 RTA00000522F.o.06.1 M00001657D:A09 26860 4012 3/24/98 413 RTA00000425F.i.10.1 M00001661D:D05 2478 4014 3/24/98 412 RTA00000425F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000425F.i.11.1 M00001669B:C12 1070 4016 3/24/98 202 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000522F.o.20.1 M00001670A:F09 75204 4018 3/24/98 331 RTA00000522F.o.20.1 M00001670A:F09 75204 4019 3/24/98 356 RTA00000522F.o.20.1 M00001670A:F09 75204 4020 3/24/98 356 RTA00000522F.o.20.1 M00001693A:H06 75804 4020 3/24/98 31	4007	3/24/98	172	RTA00000425F.f.11.1	M00001656C:C04	79275
4010 3/24/98 56 RTA00000522F.n.16.1 M00001657D:A10 26769 4011 3/24/98 20 RTA00000522F.o.06.1 M00001659D:A09 26860 4012 3/24/98 38 RTA00000528F.i.22.1 M00001661D:D05 2478 4013 3/24/98 413 RTA00000425F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000528F.j.11.1 M00001664B:F06 21716 4015 3/24/98 202 RTA00000522F.o.20.1 M00001669B:C12 1070 4016 3/24/98 432 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000522F.o.20.1 M00001670A:F09 75204 4018 3/24/98 331 RTA00000523F.a.07.1 M00001678C:F09 1981 4019 3/24/98 356 RTA000000523F.a.07.1 M00001693A:H06 75804 4020 3/24/98 14 RTA000000527F.a.13.1 M000003806C:A06 65163 4021 3/24/98 17	4008	3/24/98	58	RTA00000527F.k.06.1	M00003981B:B12	12469
4011 3/24/98 20 RTA00000522F.o.06.1 M00001659D:A09 26860 4012 3/24/98 38 RTA00000528F.i.22.1 M00001661D:D05 2478 4013 3/24/98 413 RTA00000425F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000425F.i.11.1 M00001664B:F06 21716 4015 3/24/98 402 RTA00000528F.j.11.1 M00001669B:C12 1070 4016 3/24/98 432 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 432 RTA00000522F.o.20.1 M00001670A:F09 75204 4018 3/24/98 331 RTA0000052F.o.20.1 M00001693A:H06 75804 4020 3/24/98 356 RTA00000525F.o.20.1 M00003805D:E06 37740 4021 3/24/98 14 RTA00000525F.o.21 M00003806C:A06 65163 4022 3/24/98 177 RTA00000525F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181	4009	3/24/98	240	RTA00000522F.n.14.1	M00001657C:C11	73410
4012 3/24/98 38 RTA00000528F.i.22.1 M00001661D:D05 2478 4013 3/24/98 413 RTA00000425F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000425F.i.11.1 M00001664B:F06 21716 4015 3/24/98 202 RTA00000528F.j.11.1 M00001669C:B09 74853 4016 3/24/98 432 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204 4018 3/24/98 331 RTA00000523F.b.02.1 M00001678C:F09 1981 4019 3/24/98 356 RTA00000523F.a.07.1 M00001678C:F09 1981 4020 3/24/98 356 RTA00000523F.a.07.1 M00001693A:H06 75804 4021 3/24/98 14 RTA00000523F.b.02.1 M00003805D:E06 37740 4022 3/24/98 17 RTA00000525F.n.12.1 M00003829C:H05 37575 4024 2/24/98 118	4010	3/24/98	56	RTA00000522F.n.16.1	M00001657D:A10	
4013 3/24/98 413 RTA00000425F.i.10.1 M00001664B:E08 78736 4014 3/24/98 412 RTA00000425F.i.11.1 M00001664B:F06 21716 4015 3/24/98 202 RTA00000528F.j.11.1 M00001669B:C12 1070 4016 3/24/98 432 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204 4018 3/24/98 331 RTA00000523F.a.07.1 M00001678C:F09 1981 4019 3/24/98 356 RTA00000523F.a.07.1 M00001693A:H06 75804 4020 3/24/98 200 RTA00000523F.a.07.1 M00003805D:E06 37740 4021 3/24/98 14 RTA00000525F.a.13.1 M00003806C:A06 65163 4022 3/24/98 177 RTA00000525F.n.12.1 M00001656A:H12 74117 4023 2/24/98 115 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181	4011	3/24/98	20	RTA00000522F.o.06.1	M00001659D:A09	26860
4014 3/24/98 412 RTA00000425F.i.11.1 M00001664B:F06 21716 4015 3/24/98 202 RTA00000528F.j.11.1 M00001669B:C12 1070 4016 3/24/98 432 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204 4018 3/24/98 331 RTA00000523F.a.07.1 M00001678C:F09 1981 4019 3/24/98 356 RTA00000523F.a.07.1 M00001693A:H06 75804 4020 3/24/98 200 RTA00000523F.b.02.1 M00003805D:E06 37740 4021 3/24/98 14 RTA00000523F.b.02.1 M00003806C:A06 65163 4022 3/24/98 17 RTA00000522F.n.12.1 M00001656A:H12 74117 4023 2/24/98 1158 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA0000041F.g.07.1 M00003820B:D10 21421 4026 2/24/98 127	4012	3/24/98	38	RTA00000528F.i.22.1	M00001661D:D05	2478
4015 3/24/98 202 RTA00000528F.j.11.1 M00001669B:C12 1070 4016 3/24/98 432 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204 4018 3/24/98 331 RTA00000528F.k.10.1 M00001678C:F09 1981 4019 3/24/98 356 RTA00000523F.a.07.1 M00001693A:H06 75804 4020 3/24/98 200 RTA00000527F.a.13.1 M00003805D:E06 37740 4021 3/24/98 14 RTA00000523F.b.02.1 M00003806C:A06 65163 4022 3/24/98 177 RTA00000522F.n.12.1 M00001656A:H12 74117 4023 2/24/98 1158 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA00000419F.d.07.1 M00003820B:D10 21421 4025 2/24/98 127 RTA00000411F.g.05.1 M0000382D:D06 66065 4028 2/24/98 21	4013	3/24/98	413	RTA00000425F.i.10.1	M00001664B:E08	78736
4016 3/24/98 432 RTA00000522F.o.20.1 M00001669C:B09 74853 4017 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204 4018 3/24/98 331 RTA00000528F.k.10.1 M00001678C:F09 1981 4019 3/24/98 356 RTA00000523F.a.07.1 M00001693A:H06 75804 4020 3/24/98 200 RTA00000527F.a.13.1 M00003805D:E06 37740 4021 3/24/98 14 RTA00000523F.b.02.1 M00003806C:A06 65163 4022 3/24/98 177 RTA00000522F.n.12.1 M00001656A:H12 74117 4023 2/24/98 1158 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA00000346F.f.14.1 M00003820B:D10 21421 4025 2/24/98 1227 RTA0000041F.g.05.1 M0000382D:B10 21421 4026 2/24/98 412 RTA0000041F.g.06.1 M0000382D:D04 45815 4029 2/24/98 21	4014	3/24/98	412	RTA00000425F.i.11.1	M00001664B:F06	21716
4017 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204 4018 3/24/98 331 RTA00000528F.k.10.1 M00001678C:F09 1981 4019 3/24/98 356 RTA00000523F.a.07.1 M00001693A:H06 75804 4020 3/24/98 200 RTA00000527F.a.13.1 M00003805D:E06 37740 4021 3/24/98 14 RTA00000523F.b.02.1 M00003806C:A06 65163 4022 3/24/98 177 RTA00000522F.n.12.1 M00001656A:H12 74117 4023 2/24/98 1158 RTA0000052F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA00000346F.f.14.1 M00003829C:H05 37575 4024 2/24/98 610 RTA00000411F.g.05.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.06.1 M00003822D:D06 66065 4028 2/24/98 21 RTA00000411F.g.08.1 M00003823B:F07 8188 4030 2/24/98 52	4015	3/24/98	202	RTA00000528F.j.11.1	M00001669B:C12	1070
4018 3/24/98 331 RTA00000528F.k.10.1 M00001678C:F09 1981 4019 3/24/98 356 RTA00000523F.a.07.1 M00001693A:H06 75804 4020 3/24/98 200 RTA00000527F.a.13.1 M00003805D:E06 37740 4021 3/24/98 14 RTA00000523F.b.02.1 M00003806C:A06 65163 4022 3/24/98 177 RTA00000522F.n.12.1 M00001656A:H12 74117 4023 2/24/98 1158 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA00000419F.d.07.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.05.1 M00003822D:B10 64664 4027 2/24/98 412 RTA00000411F.g.06.1 M00003822D:D04 45815 4028 2/24/98 1208 RTA0000041F.g.08.1 M00003823B:F07 8188 4030 2/24/98 502 RTA0000041F.g.08.1 M00003824C:D07 0 4031 2/24/98 528	4016	3/24/98	432			74853
4019 3/24/98 356 RTA00000523F.a.07.1 M00001693A:H06 75804 4020 3/24/98 200 RTA00000527F.a.13.1 M00003805D:E06 37740 4021 3/24/98 14 RTA00000523F.b.02.1 M00003806C:A06 65163 4022 3/24/98 177 RTA00000522F.n.12.1 M00001656A:H12 74117 4023 2/24/98 1158 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA00000346F.f.14.1 M00003820B:D10 21421 4025 2/24/98 610 RTA00000419F.d.07.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.05.1 M00003822D:B10 64664 4027 2/24/98 412 RTA00000411F.g.06.1 M00003822D:D04 45815 4028 2/24/98 1208 RTA00000411F.g.08.1 M00003823B:F07 8188 4030 2/24/98 502 RTA0000041F.g.24.1 M00003825B:B10 21503 4031 2/24/98 528 <td>4017</td> <td>3/24/98</td> <td>245</td> <td>RTA00000522F.p.09.1</td> <td>M00001670A:F09</td> <td>75204</td>	4017	3/24/98	245	RTA00000522F.p.09.1	M00001670A:F09	75204
4020 3/24/98 200 RTA00000527F.a.13.1 M00003805D:E06 37740 4021 3/24/98 14 RTA00000523F.b.02.1 M00003806C:A06 65163 4022 3/24/98 177 RTA00000522F.n.12.1 M00001656A:H12 74117 4023 2/24/98 1158 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA00000346F.f.14.1 M00003820B:D10 21421 4025 2/24/98 610 RTA00000419F.d.07.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.05.1 M00003822D:C06 66065 4027 2/24/98 412 RTA00000411F.g.08.1 M00003822D:D04 45815 4029 2/24/98 1208 RTA00000347F.e.24.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000341F.d.08.1 M00003825B:B10 21503 4031 2/24/98 528 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 </td <td>4018</td> <td>3/24/98</td> <td>331</td> <td>RTA00000528F.k.10.1</td> <td>M00001678C:F09</td> <td>1981</td>	4018	3/24/98	331	RTA00000528F.k.10.1	M00001678C:F09	1981
4021 3/24/98 14 RTA00000523F.b.02.1 M00003806C:A06 65163 4022 3/24/98 177 RTA00000522F.n.12.1 M00001656A:H12 74117 4023 2/24/98 1158 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA00000346F.f.14.1 M00003800B:F03 16998 4025 2/24/98 610 RTA00000419F.d.07.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.05.1 M00003822D:B10 64664 4027 2/24/98 412 RTA00000411F.g.06.1 M00003822D:C06 66065 4028 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 4029 2/24/98 1208 RTA00000347F.e.24.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000419F.c.16.1 M00003825B:B10 21503 4031 2/24/98 528 RTA00000419F.c.19.1 M00003829A:A08 64346 4033 2/24/98 637 <td>4019</td> <td>3/24/98</td> <td>356</td> <td>RTA00000523F.a.07.1</td> <td>M00001693A:H06</td> <td>75804</td>	4019	3/24/98	356	RTA00000523F.a.07.1	M00001693A:H06	75804
4022 3/24/98 177 RTA00000522F.n.12.1 M00001656A:H12 74117 4023 2/24/98 1158 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA00000346F.f.14.1 M00003800B:F03 16998 4025 2/24/98 610 RTA00000419F.d.07.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.05.1 M00003822D:B10 64664 4027 2/24/98 412 RTA00000411F.g.06.1 M00003822D:C06 66065 4028 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 4029 2/24/98 1208 RTA00000347F.e.24.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000341F.d.08.1 M00003824C:D07 0 4031 2/24/98 528 RTA00000419F.c.19.1 M00003825B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 81	4020	3/24/98	200	RTA00000527F.a.13.1	M00003805D:E06	37740
4023 2/24/98 1158 RTA00000405F.o.03.1 M00003829C:H05 37575 4024 2/24/98 1181 RTA00000346F.f.14.1 M00003800B:F03 16998 4025 2/24/98 610 RTA00000419F.d.07.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.05.1 M00003822D:B10 64664 4027 2/24/98 412 RTA00000411F.g.06.1 M00003822D:D06 66065 4028 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 4029 2/24/98 1208 RTA00000341F.d.08.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000341F.d.08.1 M00003824C:D07 0 4031 2/24/98 528 RTA00000419F.c.19.1 M00003829B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003829A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003819D:B01 65254 4035 2/24/98 430	4021	3/24/98	14	RTA00000523F.b.02.1	M00003806C:A06	65163
4024 2/24/98 1181 RTA00000346F.f.14.1 M00003800B:F03 16998 4025 2/24/98 610 RTA00000419F.d.07.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.05.1 M00003822D:B10 64664 4027 2/24/98 412 RTA00000411F.g.06.1 M00003822D:C06 66065 4028 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 4029 2/24/98 1208 RTA00000347F.e.24.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000341F.d.08.1 M00003824C:D07 0 4031 2/24/98 528 RTA0000045F.n.16.1 M00003825B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003830C:A03 65010 4036 2/24/98 430	4022	3/24/98	177	RTA00000522F.n.12.1	M00001656A:H12	74117
4025 2/24/98 610 RTA00000419F.d.07.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.05.1 M00003822D:B10 64664 4027 2/24/98 412 RTA00000411F.g.06.1 M00003822D:C06 66065 4028 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 4029 2/24/98 1208 RTA00000347F.e.24.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000341F.d.08.1 M00003824C:D07 0 4031 2/24/98 528 RTA00000419F.c.16.1 M00003825B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003830C:A03 65010 4036 2/24/98 430 RTA00000419F.e.02.1 M00003831C:G05 62963	4023	2/24/98	1158	RTA00000405F.o.03.1	M00003829C:H05	37575
4025 2/24/98 610 RTA00000419F.d.07.1 M00003820B:D10 21421 4026 2/24/98 1227 RTA00000411F.g.05.1 M00003822D:B10 64664 4027 2/24/98 412 RTA00000411F.g.06.1 M00003822D:C06 66065 4028 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 4029 2/24/98 1208 RTA00000347F.e.24.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000341F.d.08.1 M00003824C:D07 0 4031 2/24/98 528 RTA00000419F.c.16.1 M00003825B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003830C:A03 65010 4036 2/24/98 430 RTA00000419F.e.02.1 M00003831C:G05 62963	4024	2/24/98	1181	RTA00000346F.f.14.1	M00003800B:F03	16998
4027 2/24/98 412 RTA00000411F.g.06.1 M00003822D:C06 66065 4028 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 4029 2/24/98 1208 RTA00000347F.e.24.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000341F.d.08.1 M00003824C:D07 0 4031 2/24/98 528 RTA00000405F.n.16.1 M00003825B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003831D:B01 65254 4035 2/24/98 430 RTA00000419F.e.02.1 M00003831C:G05 62963	4025	2/24/98	610	RTA00000419F.d.07.1	M00003820B:D10	
4028 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 4029 2/24/98 1208 RTA00000347F.e.24.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000341F.d.08.1 M00003824C:D07 0 4031 2/24/98 528 RTA00000405F.n.16.1 M00003825B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003819D:B01 65254 4035 2/24/98 430 RTA00000419F.e.02.1 M00003831C:G05 62963	4026	2/24/98	1227	RTA00000411F.g.05.1	M00003822D:B10	64664
4029 2/24/98 1208 RTA00000347F.e.24.1 M00003823B:F07 8188 4030 2/24/98 502 RTA00000341F.d.08.1 M00003824C:D07 0 4031 2/24/98 528 RTA00000405F.n.16.1 M00003825B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003819D:B01 65254 4035 2/24/98 754 RTA00000419F.e.02.1 M00003830C:A03 65010 4036 2/24/98 430 RTA00000419F.e.04.1 M00003831C:G05 62963	4027	2/24/98	412	RTA00000411F.g.06.1	M00003822D:C06	66065
4030 2/24/98 502 RTA00000341F.d.08.1 M00003824C:D07 0 4031 2/24/98 528 RTA00000405F.n.16.1 M00003825B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003819D:B01 65254 4035 2/24/98 754 RTA00000419F.e.02.1 M00003830C:A03 65010 4036 2/24/98 430 RTA00000419F.e.04.1 M00003831C:G05 62963	4028	2/24/98	21	RTA00000411F.g.08.1	M00003822D:D04	45815
4031 2/24/98 528 RTA00000405F.n.16.1 M00003825B:B10 21503 4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003819D:B01 65254 4035 2/24/98 754 RTA00000419F.e.02.1 M00003830C:A03 65010 4036 2/24/98 430 RTA00000419F.e.04.1 M00003831C:G05 62963	4029	2/24/98	1208	RTA00000347F.e.24.1	M00003823B:F07	8188
4032 2/24/98 15 RTA00000419F.c.19.1 M00003820A:A08 64346 4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003819D:B01 65254 4035 2/24/98 754 RTA00000419F.e.02.1 M00003830C:A03 65010 4036 2/24/98 430 RTA00000419F.e.04.1 M00003831C:G05 62963	4030	2/24/98	502	RTA00000341F.d.08.1	M00003824C:D07	0
4033 2/24/98 637 RTA00000419F.d.14.1 M00003828A:D05 64945 4034 2/24/98 81 RTA00000419F.c.16.1 M00003819D:B01 65254 4035 2/24/98 754 RTA00000419F.e.02.1 M00003830C:A03 65010 4036 2/24/98 430 RTA00000419F.e.04.1 M00003831C:G05 62963	4031	2/24/98	528	RTA00000405F.n.16.1	M00003825B:B10	21503
4034 2/24/98 81 RTA00000419F.c.16.1 M00003819D:B01 65254 4035 2/24/98 754 RTA00000419F.e.02.1 M00003830C:A03 65010 4036 2/24/98 430 RTA00000419F.e.04.1 M00003831C:G05 62963	4032	2/24/98	15	RTA00000419F.c.19.1	M00003820A:A08	64346
4035 2/24/98 754 RTA00000419F.e.02.1 M00003830C:A03 65010 4036 2/24/98 430 RTA00000419F.e.04.1 M00003831C:G05 62963	4033	2/24/98	637	RTA00000419F.d.14.1	M00003828A:D05	64945
4036 2/24/98 430 RTA00000419F.e.04.1 M00003831C:G05 62963	4034	2/24/98	81	RTA00000419F.c.16.1	M00003819D:B01	65254
4036 2/24/98 430 RTA00000419F.e.04.1 M00003831C:G05 62963	4035	2/24/98	754	RTA00000419F.e.02.1	M00003830C:A03	65010
	4036	2/24/98	430	RTA00000419F.e.04.1		
	4037	2/24/98	541	RTA00000411F.h.15.1	M00003832A:A09	65160

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
4038	2/24/98	1079	RTA00000419F.e.10.1	M00003833B:B03	63225
4039	2/24/98	577	RTA00000419F.e.11.1	M00003833B:C12	36780
4040	2/24/98	1220	RTA00000419F.e.23.1	M00003834B:G04	65772
4041	2/24/98	691	RTA00000354R.n.08.1	M00003835A:A09	8802
4042	2/24/98	536	RTA00000411F.i.02.1	M00003835B:H11	66975
4043	2/24/98	421	RTA00000419F.f.10.1	M00003835D:G06	66193
4044	2/24/98	1150	RTA00000411F.g.24.1	M00003825B:B11	65233
4045	2/24/98	533	RTA00000423F.e.11.1	M00003809B:E10	2566
4046	2/24/98	520	RTA00000406F.c.20.1	M00003871D:G06	38578
4047	2/24/98	41	RTA00000419F.b.12.1	M00003806B:C09	63148
4048	2/24/98	917	RTA00000423F.e.21.1	M00003806B:G05	66961
4049	2/24/98	326	RTA00000419F.b.15.1	M00003806D:D11	43969
4050	2/24/98	297	RTA00000419F.b.18.1	M00003808D:D08	67034
4051	2/24/98	139	RTA00000419F.b.19.1	M00003809A:C01	65534
4052	2/24/98	1021	RTA00000419F.b.21.1	M00003809A:F01	65366
4053	2/24/98	1152	RTA00000405F.m.07.1	M00003809B:B02	37733
4054	2/24/98	310	RTA00000419F.d.06.1	M00003820B:D07	65496
4055	2/24/98	120	RTA00000401F.m.02.1	M00003920D:D07	1573
4056	2/24/98	69	RTA00000405F.o.18.1	M000033071:101 M00003839A:D07	11016
4057	2/24/98	482	RTA00000411F.e.24.1	M00003833A:B02	64781
4058	2/24/98	50	RTA00000411F.f.02.1	M00003813A:D08	63386
4059	2/24/98	602	RTA00000411F.f.06.1	M00003813A:B08	64186
4060	2/24/98	761	RTA00000411F.f.14.1	M00003813B:E09 M00003814B:C12	62984
4061	2/24/98	674	RTA00000411F.f.17.1	M00003814B:F12	65661
4062	2/24/98	1164	RTA000004111.1.17.11 RTA00000405F.m.21.1	M00003814B.F12 M00003815C:C06	24218
4063	2/24/98	951	RTA000004031.iii.21.1 RTA00000419F.c.04.1	M00003815C:C00 M00003815C:D12	63749
4064	2/24/98	471	RTA00000419F.c.11.1	M00003817B:C04	65504
4065	2/24/98	1047	RTA00000419F.c.11.1	M00003817B:C04 M00003819B:G01	65727
4066	2/24/98	1178	RTA00000419F.C.14.1 RTA00000400F.f.11.1	M00003819B:G01 M00001636A:E07	4088
4067	2/24/98	89	RTA00000400F.c.08.1	M00001030A.E07	22387
4068	2/24/98	94	RTA00000406F.a.23.1	M00003870C.A10 M00003867B:D10	38712
4069	2/24/98	1038	RTA00000406F.b.01.1	M00003867B:G07	39006
4070	2/24/98	783	RTA00000406F.b.02.1	M00003867B:G07	38744
4071	2/24/98	563	RTA00000406F.b.02.1	3.4000000000000000000000000000000000000	
4072	2/24/98	1072	RTA00000419F.j.03.1	M00003867D:A06 M00003868B:G06	18258
4072	2/24/98	846	RTA00000419F.j.03.1 RTA00000419F.j.11.1	M00003868C:C07	77578 73183
4074	2/24/98	17	RTA00000419F.j.11.1 RTA00000411F.m.15.1	M00003868D:B09	78014
4075	2/24/98	589	RTA00000411F.m.13.1	M00003868D:D09	75629
4076	2/24/98	971	RTA00000411F.i.11.1	M00003808D.D09 M00003837C:E05	
4077	2/24/98	794	RTA00000411F.I.11.1 RTA00000406F.c.06.1	M00003870C:A01	66849
4078	2/24/98	788	RTA00000400F.c.00.1 RTA00000419F.i.04.1	M00003870C:A01	37924
4078	2/24/98	883	RTA00000419F.i.04.1 RTA00000406F.c.09.1	M00003870C:E10	65791
4080	2/24/98	918	RTA00000410F.j.22.1		5671 72525
4081	2/24/98	757	•	M00003871A:A02	73525
4081	2/24/98 2/24/98	208	RTA00000423F.h.13.1	M00003871A:B09	14398
4082			RTA00000419F.j.23.1	M00003871A:C11	74470
4083	2/24/98	1127	RTA00000401F.g.22.1	M00003871A:G09	1147
4084	2/24/98	1205	RTA00000419F.k.05.1	M00003871C:E04	11757
4085	2/24/98 2/24/98	522 459	RTA00000406F.c.18.1	M00003871C:F12	14368
4080	2/24/98 2/24/98		RTA00000411E = 06.1	M00003871D:A10	78493
TU0/	4144170	965	RTA00000411F.n.06.1	M00003871D:E11	73886

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
4088	2/24/98	457	RTA00000411F.m.19.1	M00003868D:D11	74924
4089	2/24/98	145	RTA00000419F.g.12.1	M00003842C:G03	66171
4090	2/24/98	633	RTA00000341F.d.02.1	M00003797A:G03	4706
4091	2/24/98	1026	RTA00000419F.f.18.1	M00003839D:E11	64047
4092	2/24/98	524	RTA00000419F.f.23.1	M00003840D:H10	65002
4093	2/24/98	204	RTA00000351R.k.19.1	M00003841B:E03	936
4094	2/24/98	968	RTA00000419F.f.24.1	M00003841B:E06	18717
4095	2/24/98	209	RTA00000411F.j.02.1	M00003841C:D07	65310
4096	2/24/98	1118	RTA00000411F.j.03.1	M00003841C:F01	66263
4097	2/24/98	470	RTA00000411F.j.06.1	M00003841C:H08	63545
4098	2/24/98	1153	RTA00000411F.j.07.1	M00003841C:H11	66963
4099	1/28/98	412	RTA00000195AF.c.24.1	M00003860D:H07	0
4099	2/24/98	678	RTA00000195AF.c.24.1	M00003860D:H07	0
4100	2/24/98	777	RTA00000419F.g.02.1	M00003842A:A03	62839
4101	2/24/98	678	RTA00000195AF.c.24.1	M00003860D:H07	0
4101	1/28/98	412	RTA00000195AF.c.24.1	M00003860D:H07	0
4102	2/24/98	799	RTA00000411F.j.15.1	M00003843A:E04	66871
4103	2/24/98	932	RTA00000405F.p.03.1	M00003844A:A11	11346
4104	2/24/98	266	RTA00000419F.g.15.1	M00003844D:A07	32519
4105	2/24/98	547	RTA00000419F.h.02.1	M00003845D:G08	63985
4106	2/24/98	290	RTA00000411F.k.16.1	M00003852C:B06	64759
4107	2/24/98	23	RTA00000411F.k.20.1	M00003854B:A07	64973
4108	2/24/98	1138	RTA00000411F.k.21.1	M00003854B:D04	65349
4109	2/24/98	1000	RTA00000351R.j.21.1	M00003859D:C05	31604
4110	2/24/98	980	RTA00000411F.i.13.1	M00003837C:F10	66138
4111	2/24/98	112	RTA00000422F.c.11.1	M00003841D:A04	2643
4112	2/24/98	905	RTA00000405F.g.21.2	M00001673B:F07	38966
4112	2/24/98	906	RTA00000405F.g.21.1	M00001673B:F07	38966
4113	2/24/98	294	RTA00000405F.I.17.1	M00003805A:F02	17225
4114	2/24/98	105	RTA00000346F.d.08.1	M00001671A:A10	39955
4115	2/24/98	1190	RTA00000405F.g.02.2	M00001671B:G05	10567
4116	2/24/98	280	RTA00000418F.p.15.1	M00001671C:C11	31066
4117	2/24/98	1151	RTA00000405F.g.18.2	M00001672D:E08	5255
4118	2/24/98	66	RTA00000405F.g.19.2	M00001673A:G08	37150
4119	2/24/98	1239	RTA00000340F.o.22.1	M00001673B:B07	7356
4120	2/24/98	906	RTA00000405F.g.21.1	M00001673B:F07	38966
4120	2/24/98	905	RTA00000405F.g.21.2	M00001673B:F07	38966
4121	2/24/98	893	RTA00000418F.p.10.1	M00001669D:F05	75323
4122	2/24/98	906	RTA00000405F.g.21.1	M00001673B:F07	38966
4122	2/24/98	905	RTA00000405F.g.21.2	M00001673B:F07	38966
4123	2/24/98	808	RTA00000418F.p.08.1	M00001669D:D06	73983
4124	2/24/98	469	RTA00000405F.g.24.1	M00001673D:D06	39076
4125	2/24/98	1094	RTA00000405F.h.03.2	M00001673D:F10	20633
4126	2/24/98	803	RTA00000405F.h.05.2	M00001674A:G07	75706
4127	2/24/98	667	RTA00000405F.h.07.2	M00001674A:G11	4984
4128	2/24/98	276	RTA00000423F.a.18.1	M00001675A:G10	26761
4129	2/24/98	1050	RTA00000405F.f.05.2	M00001669C:D09	14359
4129	2/24/98	1049	RTA00000405F.f.05.1	M00001669C:D09	14359
4130	2/24/98	1050	RTA00000405F.f.05.2	M00001669C:D09	14359
4130	2/24/98	1049	RTA00000405F.f.05.1	M00001669C:D09	14359

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln		14000014770 404	1.600
4131	2/24/98	104	RTA00000421F.n.03.1	M00001675C:A04	1638
4132	2/24/98	388	RTA00000411F.a.07.1	M00001675C:C03	74547
4133	2/24/98	906	RTA00000405F.g.21.1	M00001673B:F07	38966
4133	2/24/98	905	RTA00000405F.g.21.2	M00001673B:F07	38966
4134	2/24/98	222	RTA00000405F.e.09.1	M00001663C:F12	38978
4135	2/24/98	518	RTA00000410F.m.18.1	M00001660B:A09	76365
4136	2/24/98	218	RTA00000346F.e.13.1	M00001660B:D03	74653
4137	2/24/98	427	RTA00000410F.m.20.1	M00001660B:E03	74285
4138	2/24/98	1099	RTA00000400F.m.16.1	M00001660B:E04	3307
4139	2/24/98	775	RTA00000405F.c.22.1	M00001660C:B06	39053
4140	2/24/98	28	RTA00000422F.p.06.2	M00001661A:B11	39282
4141	2/24/98	108	RTA00000418F.o.18.1	M00001661B:F06	78676
4142	2/24/98	954	RTA00000410F.n.05.1	M00001662A:C07	77830
4143	2/24/98	1182	RTA00000346F.d.21.1	M00001670B:G12	6641
4144	2/24/98	1043	RTA00000423F.b.17.1	M00001662B:F06	8200
4145	2/24/98	447	RTA00000423F.b.04.3	M00001675D:E10	6311
4146	2/24/98	305	RTA00000418F.p.06.1	M00001664A:F08	32628
4147	2/24/98	1116	RTA00000410F.o.04.1	M00001664D:F04	79018
4148	2/24/98	320	RTA00000422F.p.07.2	M00001661A:E06	39024
4149	2/24/98	197	RTA00000410F.o.05.1	M00001669A:B02	75262
4150	2/24/98	738	RTA00000422F.n.20.1	M00001669B:B12	38676
4151	2/24/98	495	RTA000004221.ii.20.1	M00001669C:C08	16259
4152	2/24/98	1050	RTA000004001.0.21.1 RTA00000405F.f.05.2	M00001669C:D09	14359
4152	2/24/98	1030	RTA000004051.1.05.2	M00001669C:D09	14359
4153	2/24/98	1049	RTA00000405F.f.05.1	M00001669C:D09	14359
4153	2/24/98	1049	RTA00000405F.f.05.1	M00001669C:D09	14359
4154	2/24/98	492	RTA000004031.1.03.2 RTA00000340F.o.18.1	M00001669D:C03	4261
4155	2/24/98	61	RTA00000340F.0.18.1 RTA00000410F.n.07.1	M00001662A:G01	78823
4156	2/24/98	299	RTA00000410F.II.07.1 RTA00000405F.I.15.1	M00001694A:E03	19575
4157	2/24/98	475	RTA00000403F.i.13.1 RTA00000411F.d.05.1	M00001694A.E03	75812
4157	2/24/98	692	RTA00000411F.d.03.1 RTA00000411F.d.10.1	M00001681D:C12	75612 76445
4158	2/24/98	336	RTA00000411F.d.10.1 RTA00000340F.n.13.1	M00001681D:C12	17055
4159	2/24/98	270	RTA00000340F.il.13.1 RTA00000411F.d.15.1	M00001682A:B06	74890
		969	RTA00000411F.d.13.1	M00001692A:G06	76063
4161	2/24/98		RTA00000411F.d.18.1		76063 74794
4162	2/24/98	927	RTA00000411F.d.21.1 RTA00000405F.l.03.1	M00001692B:E01 M00001692D:B01	38580
4163	2/24/98	1133	RTA00000403F.i.03.1 RTA00000401F.d.15.2	M00001692D:B01	
4164	2/24/98	576			5297
4165	2/24/98	1059	RTA00000405F.h.21.2	M00001675C:D12	39072
4166	2/24/98	780 032	RTA00000410F - 18 1	M00001693D:E08	2055
4167	2/24/98	933	RTA00000411F - 02.1	M00001680A:B02	78484
4168	2/24/98	631	RTA00000411F.e.03.1	M00001694D:C12	73648
4169	2/24/98	585	RTA00000340R.o.12.1	M00003746C:E02	53732
4170	2/24/98	604	RTA00000351R.c.13.1	M00003747D:C05	11476
4171	2/24/98	187	RTA00000351R.g.11.1	M00003779D:E08	3077
4172	2/24/98	1060	RTA00000346F.g.02.1	M00003792A:B10	6901
4173	2/24/98	690	RTA00000341F.b.05.1	M00003793D:A11	0
4174	2/24/98	86	RTA00000346F.g.22.1	M00003794D:G03	6371
4175	2/24/98	1051	RTA00000346F.h.24.1	M00003797A:C11	4379
4176	2/24/98	377	RTA00000346F.i.01.1	M00003797A:D06	22260
4177	2/24/98	963	RTA00000405F.I.07.1	M00001693C:E09	38636

Appln	SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
4179 2/24/98 781		Appln	Appln			
A180 2/24/98 82						78544
4181 2/24/98 465				RTA00000423F.f.09.1	M00003808C:A05	64823
4182 2/24/98 465	4180			RTA00000346F.d.12.1	M00001676B:B09	11777
4183				RTA00000411F.b.03.1	M00001676B:E01	23634
4184 2/24/98 789	4182	2/24/98	465	RTA00000350R.p.18.1	M00001676B:F05	11460
4185 2/24/98 267 RTA00000423F.a.19.1 M00001676D:A02 21396 4186 2/24/98 836 RTA0000041F.b.17.1 M0000167D:B02 72893 4187 2/24/98 370 RTA0000045F.i.20.1 M00001677B:G11 38532 4188 2/24/98 39 RTA00000187AF.i.7.1 M0000167B:B04 72852 4189 2/24/98 389 RTA00000195AF.c.2.1 M0000167B:B04 72852 4191 2/24/98 958 RTA00000195AF.c.8.1 M0000167B:B101 0 4191 1/28/98 520 RTA00000195AF.c.8.1 M0000167B:H01 0 4192 1/28/98 520 RTA00000195AF.c.8.1 M0000167B:H01 0 4193 1/28/98 520 RTA00000149C:F.c.8.1 M0000167B:H01 0 4194 2/24/98 500 RTA00000340F.n.01.1 M00001679A:G06 39081 4195 2/24/98 337 RTA00000340F.n.01.1 M00001679A:G06 39081 4196 2/24/98 337 RTA000004	4183	2/24/98	56	RTA00000411F.b.06.1	M00001676C:A04	77884
4186 2/24/98 836 RTA0000041 F.b.17.1 M00001676D:B02 728/93 4187 2/24/98 370 RTA00000187AF.I.20.1 M00001677A:G11 38532 4188 2/24/98 389 RTA00000187AF.I.7.1 M00001677B:B04 72852 4190 2/24/98 389 RTA00000187AF.I.2.1 M0000167B:B04 72852 4190 2/24/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 2/24/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 2/24/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4193 2/24/98 500 RTA00000195AF.c.8.1 M00001678B:H01 0 4194 2/24/98 323 RTA00000411F.c.10.1 M00001678D:B11 73117 4194 2/24/98 323 RTA00000340F.n.01.1 M00001679A:D10 16409 4195 2/24/98 337 RTA00000340F.n.01.1 M00001679D:B02 78533 4197 1/28/98 238 RTA00000340F.n.01.1 M00001679D:F02 78415 4197 2/24/98 407 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000187AR.k.12.1 M00001679D:F02 78415 4198 2/24/98 407 RTA00000340R.m.07.1 M00001679D:F02 78415 4199 2/24/98 48 RTA00000141F.a.15.1 M00001679D:F02 78415 4199 2/24/98 48 RTA00000141F.a.15.1 M00001679D:F02 78415 4200 2/24/98 48 RTA00000141F.a.15.1 M00001679D:F02 78415 4201 2/24/98 48 RTA00000141F.a.15.1 M00001679D:F02 78415 4201 2/24/98 48 RTA00000195AF.d.4.1 M000038781D:D06 22766 4201 1/28/98 138 RTA0000041F.a.15.1 M00003878C:F06 22961 4204 2/24/98 300 RTA00000406F.f.05.1 M00003878C:F06 22961 4204 2/24/98 316 RTA00000406F.f.05.1 M00003878C:F06 22961 4204 2/24/98 36 RTA00000406F.f.05.1 M00003878C:F06 22961 4204 2/24/98 36 RTA00000406F.f.05.1 M00003878C:F06 22961 4204 2/24/98 36 RTA00000406F.f.05.1 M00003878C:G02 5040 4210 2/24/98 36 RTA00000406F.f.05.1 M00003879C:D02 37972 4213 2/24/98 36 RTA00000406F.f.05.1 M00003879C:D02 37972 4213 2/24/98 36 RTA00000406F.f.05.1	4184	2/24/98	789	RTA00000423F.b.13.1	M00001676C:E07	20619
4187 2/24/98 370	4185	2/24/98	267	RTA00000423F.a.19.1	M00001676D:A02	21396
4188 2/24/98 39 RTA00000187AF.I.7.1 M00001680D:F08 10539 4189 2/24/98 389 RTA00000411F.c.02.1 M0000167B:B04 72852 4190 2/24/98 1004 RTA00000195AF.c.8.1 M0000167B:B101 0 4191 1/28/98 520 RTA00000195AF.c.8.1 M0000167B:H01 0 4192 1/28/98 520 RTA00000195AF.c.8.1 M0000167B:H01 0 4192 1/28/98 520 RTA00000195AF.c.8.1 M0000167B:H01 0 4193 2/24/98 520 RTA0000041F.c.10.1 M0000167B:H01 0 4194 1/28/98 520 RTA00000421F.n.19.1 M0000167B:B10 0 4195 2/24/98 309 RTA00000340F.n.01.1 M00001679A:D10 16409 4195 2/24/98 337 RTA00000340F.n.01.1 M00001679D:B02 78533 4197 1/28/98 238 RTA00000340F.n.01.1 M00001679D:B02 78415 4198 1/28/98 407 RTA00000187AR.k.1	4186	2/24/98	836	RTA00000411F.b.17.1	M00001676D:B02	72893
4188 2/24/98 39 RTA00000187AF.I.7.1 M00001680D:F08 10539 4189 2/24/98 389 RTA00000411F.c.02.1 M00001677B:B04 72852 4190 2/24/98 958 RTA00000195AF.c.8.1 M00001678B:H01 0 4191 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 2/24/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4193 2/24/98 520 RTA00000340F.n.01.1 M00001679A:D10 16409 4194 2/24/98 323 RTA00000340F.n.01.1 M00001679A:D10 16409 4195 2/24/98 337 RTA00000340F.n.04.1 M00001679D:B02 78533 4197 1/28/98 238 RTA00000187AR.k.12.1 M00001679D:F02 78415 4198 1/28/98 407 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 2/24/98 407 RT	4187	2/24/98	370	RTA00000405F.i.20.1	M00001677A:G11	38532
4190 2/24/98 1004 RTA00000419F.a.24.1 M00001680B:D02 79290 4191 2/24/98 958 RTA00000195AF.c.8.1 M00001678B:H01 0 4191 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 2/24/98 958 RTA00000195AF.c.8.1 M00001678B:H01 0 4193 2/24/98 500 RTA00000411F.c.10.1 M00001678B:H01 0 4194 2/24/98 500 RTA00000411F.c.10.1 M00001679A:D10 16409 4195 2/24/98 323 RTA00000340F.n.01.1 M00001679A:G06 39081 4196 2/24/98 337 RTA00000340F.n.01.1 M00001679D:B02 78533 4197 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4199 2/24/98 407 RTA000004340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 387	4188	2/24/98	39	RTA00000187AF.I.7.1		
4190 2/24/98 1004 RTA00000419F.a.24.1 M00001680B:D02 79290 4191 2/24/98 958 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 1/28/98 958 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4193 2/24/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4193 2/24/98 520 RTA00000411F.c.10.1 M00001678B:H01 0 4194 2/24/98 323 RTA00000340F.n.01.1 M00001679A:D10 16409 4195 2/24/98 337 RTA00000340F.n.01.1 M00001679D:B02 78533 4197 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 387 RTA000004340R.m.07.1 M00001679D:F02 78415 4199 2/24/98 48 RTA00	4189	2/24/98	389	RTA00000411F.c.02.1	M00001677B:B04	72852
4191 2/24/98 958 RTA00000195AF.c.8.1 M00001678B:H01 0 4191 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 2/24/98 958 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4193 2/24/98 500 RTA0000041F.c.8.1 M00001678B:H01 0 4194 2/24/98 323 RTA00000421F.n.19.1 M00001679A:D10 16409 4195 2/24/98 339 RTA00000340F.n.01.1 M00001679A:G06 39081 4196 2/24/98 337 RTA00000340F.p.04.1 M00001679D:F02 78415 4197 1/28/98 238 RTA00000187AR.k.12.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 387 RTA0000041F.a.12.1 M00001679D:F02 78415 4198 1/28/98 48 RTA000004	4190	2/24/98	1004	RTA00000419F.a.24.1	M00001680B:D02	79290
4191 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 2/24/98 958 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4193 2/24/98 500 RTA0000041F.n.l0.1 M00001679A:D10 16409 4194 2/24/98 323 RTA00000340F.n.l0.1 M00001679A:G06 39081 4196 2/24/98 337 RTA00000340F.n.l0.1 M00001679D:F02 78533 4197 1/28/98 238 RTA00000187AR.k.l2.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000043PAR.k.l2.1 M00001679D:F02 78415 4199 2/24/98 387 RTA0000041F.a.l5.1 M00001679D:F02 78415 4200 2/24/98 48 <t< td=""><td>4191</td><td>2/24/98</td><td>958</td><td>RTA00000195AF.c.8.1</td><td></td><td>0</td></t<>	4191	2/24/98	958	RTA00000195AF.c.8.1		0
4192 2/24/98 958 RTA00000195AF.c.8.1 M00001678B:H01 0 4192 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4193 2/24/98 500 RTA0000041IF.c.10.1 M00001678B:H01 73117 4194 2/24/98 323 RTA00000340F.n.01.1 M00001679A:G06 39081 4195 2/24/98 337 RTA00000340F.p.04.1 M00001679D:B02 78533 4197 1/28/98 238 RTA00000340F.p.04.1 M00001679D:F02 78415 4197 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 387 RTA0000041F.a.15.1 M00001679D:F02 78415 4199 2/24/98 387 RTA0000041F.a.15.1 M00001677B:A12 30041 4200 2/24/98 387 RTA0000041F.a.15.4.1 M00001677B:A12 30041 4201 1/28/98 185	4191	1/28/98	520	RTA00000195AF.c.8.1		
4192 1/28/98 520 RTA00000195AF.c.8.1 M00001678B:H01 0 4193 2/24/98 500 RTA0000041IF.c.10.1 M00001678D:B11 73117 4194 2/24/98 323 RTA00000340F.n.01.1 M00001679A:G06 39081 4195 2/24/98 337 RTA00000340F.p.04.1 M00001679A:G06 39081 4196 2/24/98 337 RTA00000340F.p.04.1 M00001679D:F02 78415 4197 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000187AR.k.12.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000187AR.k.12.1 M00001679D:F02 78415 4198 1/28/98 337 RTA00000411F.a.15.1 M00001679D:F02 78415 4199 2/24/98 387 RTA00000411F.a.15.1 M00001677D:F02 78415 4199 2/24/98 387 RTA000000411F.a.15.1 M00001677B:A.12 30041 4201 1/28/98 185 <td>4192</td> <td>2/24/98</td> <td>958</td> <td></td> <td></td> <td></td>	4192	2/24/98	958			
4193 2/24/98 500 RTA00000411F.c.10.1 M00001678D:B11 73117 4194 2/24/98 323 RTA00000421F.n.19.1 M00001679A:D10 16409 4195 2/24/98 339 RTA00000340F.n.01.1 M00001679D:B02 78533 4196 2/24/98 337 RTA00000340F.n.04.1 M00001679D:F02 78513 4197 1/28/98 238 RTA00000340R.m.07.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000187AR.k.12.1 M00001679D:F02 78415 4198 1/28/98 238 RTA00000187AR.k.12.1 M00001679D:F02 78415 4198 2/24/98 407 RTA00000187AR.k.12.1 M00001679D:F02 78415 4199 2/24/98 387 RTA0000040R.m.07.1 M00001679D:F02 78415 4199 2/24/98 387 RTA00000041F.a.15.1 M00001679D:F02 78415 4199 2/24/98 387 RTA00000041F.a.15.1 M00001679D:F02 78415 4200 2/24/98 185 </td <td>4192</td> <td>1/28/98</td> <td>520</td> <td></td> <td></td> <td></td>	4192	1/28/98	520			
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4210 2/24/98 494 RTA00000419F.k.12.1 M00003876C:F02 0 4211 2/24/98 515 RTA00000419F.l.03.1 M00003879A:D02 79060 4212 2/24/98 26 RTA00000423F.h.18.1 M00003876C:D02 37972 4213 2/24/98 49 RTA00000406F.d.12.1 M00003875C:A01 38575 4214 2/24/98 986 RTA00000406F.d.24.1 M00003876B:C05 37997 4215 2/24/98 150 RTA00000419F.k.19.1 M00003877C:G12 75447 4216 2/24/98 538 RTA00000423F.g.04.1 M00003903D:C12 23012 4217 2/24/98 1046 RTA00000346F.j.06.1 M00003879A:A02 5767 4218 2/24/98 868 RTA00000406F.i.08.1 M00003879A:B08 38601 4219 2/24/98 409 RTA00000354R.m.02.1 M00003879A:B08 38601 4220 2/24/98 543 RTA00000419F.k.24.1 M00003878C:G08 75596	4209	2/24/98	1207			
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4212 2/24/98 26 RTA00000423F.h.18.1 M00003876C:D02 37972 4213 2/24/98 49 RTA00000406F.d.12.1 M00003875C:A01 38575 4214 2/24/98 986 RTA00000406F.d.24.1 M00003876B:C05 37997 4215 2/24/98 150 RTA00000419F.k.19.1 M00003877C:G12 75447 4216 2/24/98 538 RTA00000423F.g.04.1 M00003903D:C12 23012 4217 2/24/98 1046 RTA00000346F.j.06.1 M00003879A:A02 5767 4218 2/24/98 868 RTA00000406F.i.08.1 M00003879A:B08 38601 4219 2/24/98 409 RTA00000354R.m.02.1 M00003890B:C08 12766 4220 2/24/98 543 RTA00000419F.k.24.1 M00003878C:G08 75596	4211	2/24/98	515	RTA00000419F.l.03.1		
4213 2/24/98 49 RTA00000406F.d.12.1 M00003875C:A01 38575 4214 2/24/98 986 RTA00000406F.d.24.1 M00003876B:C05 37997 4215 2/24/98 150 RTA00000419F.k.19.1 M00003877C:G12 75447 4216 2/24/98 538 RTA00000423F.g.04.1 M00003903D:C12 23012 4217 2/24/98 1046 RTA00000346F.j.06.1 M00003879A:A02 5767 4218 2/24/98 868 RTA00000406F.i.08.1 M00003879A:B08 38601 4219 2/24/98 409 RTA00000406F.f.11.1 M00003879A:B08 38601 4220 2/24/98 924 RTA00000354R.m.02.1 M00003878C:G08 75596	4212	2/24/98	26	RTA00000423F.h.18.1		
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4216 2/24/98 538 RTA00000423F.g.04.1 M00003903D:C12 23012 4217 2/24/98 1046 RTA00000346F.j.06.1 M00003879A:A02 5767 4218 2/24/98 868 RTA00000406F.i.08.1 M00003903C:E12 37946 4219 2/24/98 409 RTA00000406F.f.11.1 M00003879A:B08 38601 4220 2/24/98 924 RTA00000354R.m.02.1 M00003890B:C08 12766 4221 2/24/98 543 RTA00000419F.k.24.1 M00003878C:G08 75596	4215	2/24/98	150	RTA00000419F.k.19.1	M00003877C:G12	
4217 2/24/98 1046 RTA00000346F.j.06.1 M00003879A:A02 5767 4218 2/24/98 868 RTA00000406F.i.08.1 M00003903C:E12 37946 4219 2/24/98 409 RTA00000406F.f.11.1 M00003879A:B08 38601 4220 2/24/98 924 RTA00000354R.m.02.1 M00003890B:C08 12766 4221 2/24/98 543 RTA00000419F.k.24.1 M00003878C:G08 75596	4216	2/24/98	538	RTA00000423F.g.04.1		
4218 2/24/98 868 RTA00000406F.i.08.1 M00003903C:E12 37946 4219 2/24/98 409 RTA00000406F.f.11.1 M00003879A:B08 38601 4220 2/24/98 924 RTA00000354R.m.02.1 M00003890B:C08 12766 4221 2/24/98 543 RTA00000419F.k.24.1 M00003878C:G08 75596	4217	2/24/98	1046			
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4221 2/24/98 543 RTA00000419F.k.24.1 M00003878C:G08 75596	4220	2/24/98				
		2/24/98	543			
	4222	1/28/98	185	RTA00000195AF.d.4.1		

Priority	SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
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4223 2/24/98 582 RTA0000034IF.h.10.1 M00003901B:G11 0 4224 2/24/98 550 RTA00000406F.i.13.1 M00003875C:A09 75816 4225 2/24/98 13 RTA00000406F.i.13.1 M00003904A:C04 37904 4226 2/24/98 13 RTA00000405F.i.18.1 M00003903D:D10 799 4228 2/24/98 185 RTA00000415F.i.19.1 M00003903D:D110 799 4229 2/24/98 185 RTA0000041F.i.19.1 M00003903D:H11 39080 4230 2/24/98 802 RTA0000041F.n.11. M00003875A:B01 7276 4231 2/24/98 498 RTA0000041F.n.09.1 M00003875A:B01 73963 4233 2/24/98 498 RTA0000041F.n.09.1 M00003875A:A07 78962 4234 2/24/98 984 RTA0000046F.g.08.1 M00003875R:B01 73963 4235 2/24/98 818 RTA0000046F.g.08.1 M00003875R:B11 37963 4235 2/24/98 313 RTA						22566
4224 2/24/98 550 RTA00000411F.n.20.1 M00003875C:A09 75816 4225 2/24/98 614 RTA00000406F.i.13.1 M00003904A:C04 37904 4226 2/24/98 13 RTA00000406F.i.13.1 M00003879B:G02 37598 4227 2/24/98 185 RTA0000040F.i.12.1 M00003903D:D10 799 4228 2/24/98 177 RTA00000406F.i.12.1 M00003903D:D11 39080 4230 2/24/98 802 RTA0000041F.n.11.1 M00003875A:B01 77276 4231 2/24/98 498 RTA00000406F.e.15.1 M00003875A:B01 77276 4232 2/24/98 498 RTA00000406F.e.15.1 M00003875A:A07 78962 4233 2/24/98 498 RTA00000406F.e.15.1 M00003875A:A07 78962 4234 2/24/98 84 RTA00000406F.e.05.1 M00003875A:A07 78962 4234 2/24/98 818 RTA00000406F.e.05.1 M00003877B:H10 750 4235 2/24/98 313						
4225 2/24/98 614 RTA00000406F.i.13.1 M00003904A:C04 37904 4226 2/24/98 13 RTA00000406F.f.18.1 M00003879B:G02 38587 4227 2/24/98 1256 RTA00000401F.k.19.1 M000039903C:C05 37958 4228 2/24/98 185 RTA00000406F.i.12.1 M00003903C:C05 37958 4230 2/24/98 802 RTA00000406F.g.03.1 M00003880B:D11 39080 4231 2/24/98 34 RTA0000041F.n.11 M00003875A:B01 77276 4232 2/24/98 34 RTA0000041F.n.19.1 M00003875A:B01 77276 4233 2/24/98 392 RTA0000041F.n.09.1 M00003875A:A07 78962 4234 2/24/98 84 RTA000004406F.g.08.1 M00003880C:H03 37963 4235 2/24/98 818 RTA000004406F.g.07.1 M00003880C:H03 37864 4236 2/24/98 313 RTA00000446F.g.07.1 M00003880C:H13 37925 4238 1/24/98 324						-
4226 2/24/98 13 RTA00000406F.f.18.1 M00003879B:G02 38587 4227 2/24/98 1256 RTA00000401F.k.19.1 M00003903D:D10 799 4228 2/24/98 185 RTA0000041F.k.19.1 M00003903D:D11 39080 4229 2/24/98 817 RTA0000041F.k.19.1 M00003903D:H11 39080 4230 2/24/98 802 RTA0000041F.n.11.1 M00003875A:B01 77276 4231 2/24/98 498 RTA00000406F.e.15.1 M000038775:A07 78962 4232 2/24/98 498 RTA00000406F.e.15.1 M00003875A:A07 78962 4233 2/24/98 848 RTA00000406F.g.08.1 M000038808:D10 39074 4234 2/24/98 818 RTA00000406F.g.08.1 M00003875A:A07 78962 4235 2/24/98 818 RTA00000406F.g.05.1 M0000387TB:H10 750 4237 2/24/98 313 RTA00000426F.g.07.1 M0000387TB:H10 750 4239 2/24/98 773						
4227 2/24/98 1256 RTA00000401F.k.19.1 M00003903D:D10 799 4228 2/24/98 185 RTA00000425F.j.05.1 M00003903C:D5 37958 4229 2/24/98 802 RTA00000406F.g.03.1 M00003903D:H11 39080 4230 2/24/98 802 RTA00000406F.g.03.1 M00003875C:Al1 38690 4231 2/24/98 34 RTA00000406F.g.03.1 M00003877C:Al1 39074 4232 2/24/98 498 RTA00000406F.g.01.1 M00003877C:Al1 39074 4233 2/24/98 929 RTA00000406F.g.08.1 M00003887C:H03 37963 4234 2/24/98 984 RTA00000406F.g.07.1 M00003887B:H10 37963 4235 2/24/98 592 RTA0000042F.p.91.8.1 M00003880C:El1 37963 4236 2/24/98 313 RTA0000046F.g.07.1 M00003880C:El1 37925 4238 1/28/98 324 RTA0000046F.h.03.1 M0000330B:A09 38585 4240 3/24/98 218						
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4236 2/24/98 592 RTA00000421F.p.18.1 M00003877B:H10 750 4237 2/24/98 313 RTA00000406F.g.07.1 M00003880C:E11 37925 4238 1/28/98 324 RTA00000184F.j.06.1 M00001556B:G0 11294 4239 2/24/98 773 RTA00000406F.p.09.1 M00004031B:A09 38585 4240 3/24/98 244 RTA00000426F.p.09.1 M00004033D:B07 66665 4241 1/28/98 181 RTA00000197AF.n.2.1 M00001585A:D02 6229 4244 1/28/98 877 RTA0000041F.a.09.1 M00001675C:F01 78629 4244 2/24/98 844 RTA0000041F.a.09.1 M00001675C:F01 78629 4245 2/24/98 352 RTA00000426F.m.02.1 M00004034C:C06 66237 4245 2/24/98 429 RTA00000525F.a.14.1 M00004037C:D07 7566 4248 2/24/98 118 RTA0000040F.m.13.1 M00001619B:C02 37566 4248 2/24/98 156	4234	2/24/98	984			
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4238 1/28/98 324 RTA00000184F.j.06.1 M00001556B:G02 11294 4239 2/24/98 773 RTA00000406F.h.03.1 M00003901B:A09 38585 4240 3/24/98 244 RTA00000426F.p.09.1 M00004033D:B07 66665 4241 3/24/98 222 RTA00000426F.p.10.1 M00004033D:C05 65845 4242 1/28/98 181 RTA00000198AF.d.2.1 M00001585A:F07 0 4243 1/28/98 77 RTA00000197AF.n.2.1 M00001535A:D02 6229 4244 2/24/98 352 RTA00000411F.a.09.1 M00001675C:G01 73073 4246 3/24/98 272 RTA00000426F.m.02.1 M00001675C:G01 73073 4246 3/24/98 429 RTA00000426F.m.02.1 M00001675C:G01 73073 4247 3/24/98 429 RTA00000426F.m.03.1 M00001479D:H003 78382 4249 3/24/98 118 RTA00000409F.m.13.1 M00001437C:D07 16679 4250 2/24/98 1198 RTA00000404F.h.20.1 M00001618B:E05 0 4251 2/24/98 1198 RTA00000412F.f.10.2 M00003959A:A03 65405 4252 2/24/98 1199 RTA00000525F.a.22.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000525F.a.22.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000404F.h.20.1 M00001619B:A09 15564 4255 2/24/98 119 RTA00000403F.g.03.1 M00001619B:A09 15564 4255 2/24/98 562 RTA00000525F.a.22.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000403F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA00000403F.a.24.1 M00004037B:A04 24715 4258 3/24/98 43 RTA00000418F.g.03.1 M00001479D:G06 23537 4266 2/24/98 53 RTA00000409F.m.02.1 M000004029B:H08 34149 4258 3/24/98 43 RTA00000409F.m.02.1 M000004029B:H08 34149 4258 3/24/98 43 RTA00000409F.m.02.1 M000001616C:A02 73174 4264 3/24/98 1201 RTA00000409F.m.02.1 M00001616C:A02 73174 4264 3/24/98 131 RTA00000409F.m.02.1 M00001616C:A02 73174 4264 2/24/98 131 RTA00000409F.m.02.1 M00001616C:A02 73174 4266 1/28/98 131 RTA00000409F.m.02.1 M00001619D:C03 6539 4266 1/28/98 131 RTA00000185AR.d.11.1 M00001528A:A01 5175 4266 1/28/98 136 RTA00000183AF.h.11.1 M00001528A:A01 5175 4266 1/28/98 136 RTA00000183AF.h.11.1 M00001528A:A01 5175 4266 1/28/98 136 RTA00000183AF.h.19.1 M00001528A:A01 5175 4266 1/28/98 136 RTA00000183AF.h.19.1 M00001528A:A01 5175 4266 1/28/98 236 RTA00000183AF.h.19.1 M000001528A:A01 5175 4266 1/28/98 236 RTA00000527F.o.01.1 M000004027A:D06 19088	4236	2/24/98	592	RTA00000421F.p.18.1	M00003877B:H10	750
4239 2/24/98 773 RTA00000406F.h.03.1 M00003901B:A09 38585 4240 3/24/98 244 RTA00000426F.p.09.1 M00004033D:B07 66665 4241 3/24/98 222 RTA00000198AF.d.2.1 M00001585A:F07 0 4242 1/28/98 181 RTA00000197AF.n.2.1 M00001585A:F07 0 4243 1/28/98 77 RTA00000197AF.n.2.1 M00001535A:D02 6229 4244 2/24/98 844 RTA00000411F.a.09.1 M00001675C:F01 78629 4245 2/24/98 352 RTA00000426F.m.02.1 M00004038:C02 37566 4246 3/24/98 272 RTA00000525F.a.14.1 M00004038:C02 37566 4247 3/24/98 429 RTA00000426F.m.03.1 M00004037C:D07 16679 4248 2/24/98 118 RTA000000525F.a.12.1 M00004037C:D07 16679 4250 2/24/98 1198 RTA000000409F.m.13.1 M00004037C:D07 16679 4251 2/24/98 1139	4237	2/24/98	313	RTA00000406F.g.07.1	M00003880C:E11	37925
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4240 3/24/98 244 RTA00000426F.p.09.1 M00004033D:B07 66665 4241 3/24/98 222 RTA00000426F.p.10.1 M00004033D:C05 65845 4242 1/28/98 181 RTA00000197AF.n.2.1 M00001535A:D02 6229 4243 1/28/98 77 RTA00000197AF.n.2.1 M00001535A:D02 6229 4244 2/24/98 844 RTA00000411F.a.09.1 M00001675C:G01 73073 4246 3/24/98 272 RTA00000426F.m.02.1 M00004034C:C06 66237 4247 3/24/98 429 RTA00000525F.a.14.1 M00004037C:D07 16679 4248 2/24/98 118 RTA00000525F.b.22.1 M0000437C:D07 16679 4250 2/24/98 156 RTA000000525F.b.22.1 M0000437C:D07 16679 4251 2/24/98 198 RTA000000409F.m.13.1 M00001618B:E05 0 4251 2/24/98 1198 RTA00000404F.b.20.1 M000016198.A09 15564 4252 2/24/98 119	4239	2/24/98	773	RTA00000406F.h.03.1	M00003901B:A09	38585
4242 1/28/98 181 RTA00000198AF.d.2.1 M00001585A:F07 0 4243 1/28/98 77 RTA00000197AF.n.2.1 M00001535A:D02 6229 4244 2/24/98 844 RTA00000411F.a.09.1 M00001675C:F01 78629 4245 2/24/98 352 RTA00000411F.a.10.1 M00001675C:G01 73073 4246 3/24/98 272 RTA00000426F.m.02.1 M00004034C:C06 66237 4247 3/24/98 429 RTA00000525F.a.14.1 M00004033B:C02 37566 4248 2/24/98 118 RTA00000409F.m.03.1 M00001479D:H03 78382 4249 3/24/98 156 RTA00000525F.b.22.1 M00000437C:D07 16679 4250 2/24/98 70 RTA00000409F.m.13.1 M00001618B:E05 0 4251 2/24/98 1198 RTA00000412F.f.10.2 M00003959A:A03 65405 4252 2/24/98 1139 RTA00000404F.h.20.1 M00001619B:A09 15564 4253 3/24/98 41 RTA0000044F.h.20.1 M00001619B:A09 15564 4253 3/24/98 452 RTA00000525F.a.22.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000525F.a.22.1 M00000437B:A04 24715 4255 2/24/98 1019 RTA00000403F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA0000043F.a.24.1 M00001479D:G06 23537 4256 2/24/98 532 RTA0000043F.a.24.1 M00001479D:G06 23537 4257 3/24/98 43 RTA0000043F.a.24.1 M00001429B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029B:H08 34149 4259 2/24/98 303 RTA00000527F.p.07.1 M000004029B:H08 34149 4260 2/24/98 101 RTA0000049F.m.02.1 M0000161C:A11 9157 4261 2/24/98 1170 RTA0000049F.m.02.1 M0000161C:A02 73174 4264 2/24/98 176 RTA0000049F.n.02.1 M0000161C:A02 73174 4264 2/24/98 176 RTA0000049F.l.24.1 M000011579D:C03 6539 4265 1/28/98 131 RTA00000185AF.d.11.1 M00001579D:C03 6539 4266 1/28/98 131 RTA00000185AF.d.11.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000185AF.d.11.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000185AF.d.11.1 M00001528A:A01 5175 4266 1/28/98 176 RTA0000013AA.c.7.1 M00001528A:A01 5175 4266 3/24/98 90 RTA000001525F.a.03.1 M00001528A:A01 5175 4266 3/24/98 90 RTA00000525F.a.03.1 M00001528A:A01 5175 4268 3/24/98 90 RTA00000525F.a.03.1 M00001528A:A01 5175	4240	3/24/98	244	RTA00000426F.p.09.1	M00004033D:B07	66665
4242 1/28/98 181 RTA00000198AF.d.2.1 M00001585A:F07 0 4243 1/28/98 77 RTA00000197AF.n.2.1 M00001675C:F01 78629 4244 2/24/98 844 RTA000000411F.a.09.1 M00001675C:F01 78629 4245 2/24/98 352 RTA00000426F.m.02.1 M00004034C:C06 66237 4247 3/24/98 429 RTA00000525F.a.14.1 M00004033B:C02 37566 4248 2/24/98 118 RTA0000049F.h.03.1 M00001479D:H03 78382 4249 3/24/98 156 RTA00000525F.b.22.1 M00004037C:D07 16679 4250 2/24/98 1198 RTA00000449F.m.13.1 M00001618B:E05 0 4251 2/24/98 1198 RTA00000449F.h.20.1 M00003959A:A03 65405 4252 2/24/98 118 RTA0000044F.h.20.1 M00001619B:A09 15564 4253 3/24/98 41 RTA0000042F.b.12.1 M00001619B:A09 15564 4253 3/24/98 452	4241	3/24/98	222	RTA00000426F.p.10.1	M00004033D:C05	65845
4243 1/28/98 77 RTA00000197AF.n.2.1 M00001535A:D02 6229 4244 2/24/98 844 RTA00000411F.a.09.1 M00001675C:F01 78629 4245 2/24/98 352 RTA00000041F.a.10.1 M00001675C:G01 73073 4246 3/24/98 272 RTA000000525F.a.14.1 M00004034C:C06 66237 4247 3/24/98 429 RTA00000525F.a.14.1 M00004033B:C02 37566 4248 2/24/98 118 RTA000000525F.b.22.1 M00004037C:D07 16679 4250 2/24/98 156 RTA000000409F.m.13.1 M00001618B:E05 0 4251 2/24/98 1198 RTA000000412F.f.10.2 M00003959A:A03 65405 4252 2/24/98 1139 RTA000000412F.f.10.2 M00001619B:A09 15564 4253 3/24/98 41 RTA000000525F.b.17.1 M00001619B:A09 15564 4253 3/24/98 452 RTA000000525F.a.22.1 M00004037B:A04 24715 4254 3/24/98 452 </td <td></td> <td>1/28/98</td> <td>181</td> <td><u>-</u></td> <td>M00001585A:F07</td> <td>0</td>		1/28/98	181	<u>-</u>	M00001585A:F07	0
4244 2/24/98 844 RTA00000411F.a.09.1 M00001675C:F01 78629 4245 2/24/98 352 RTA00000411F.a.10.1 M00001675C:G01 73073 4246 3/24/98 272 RTA00000426F.m.02.1 M00004033B:C02 37566 4247 3/24/98 429 RTA00000525F.a.14.1 M00004037C:D07 78382 4248 2/24/98 118 RTA0000049F.h.03.1 M00004037C:D07 16679 4250 2/24/98 156 RTA00000409F.m.13.1 M00004037C:D07 16679 4251 2/24/98 1198 RTA00000409F.m.13.1 M00001618B:E05 0 4251 2/24/98 1198 RTA00000404F.h.20.1 M00001619B:A09 15564 4252 2/24/98 1139 RTA000000525F.b.17.1 M00004037B:A04 24715 4253 3/24/98 41 RTA000000525F.b.17.1 M00004037B:A04 24715 4254 3/24/98 41 RTA00000043F.g.03.1 M00001479D:G06 23537 4256 2/24/98 52				RTA00000197AF.n.2.1	M00001535A:D02	6229
4245 2/24/98 352 RTA00000411F.a.10.1 M00001675C:G01 73073 4246 3/24/98 272 RTA00000426F.m.02.1 M00004034C:C06 66237 4247 3/24/98 429 RTA00000525F.a.14.1 M00001479D:H03 78382 4248 2/24/98 118 RTA00000408F.h.03.1 M00001479D:H03 78382 4249 3/24/98 156 RTA00000525F.b.22.1 M00004037C:D07 16679 4250 2/24/98 1198 RTA00000409F.m.13.1 M00001618B:E05 0 4251 2/24/98 1198 RTA00000412F.f.10.2 M00003959A:A03 65405 4252 2/24/98 1139 RTA00000412F.f.10.2 M00001619B:A09 15564 4253 3/24/98 41 RTA00000525F.b.17.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000525F.a.22.1 M00004037B:A04 24715 4255 2/24/98 1019 RTA00000403F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532		2/24/98			M00001675C:F01	78629
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4248 2/24/98 118 RTA00000408F.h.03.1 M00001479D:H03 78382 4249 3/24/98 156 RTA00000525F.b.22.1 M00004037C:D07 16679 4250 2/24/98 70 RTA00000409F.m.13.1 M00001618B:E05 0 4251 2/24/98 1198 RTA00000412F.f.10.2 M00003959A:A03 65405 4252 2/24/98 1139 RTA00000404F.h.20.1 M00001619B:A09 15564 4253 3/24/98 41 RTA00000525F.b.17.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000525F.a.22.1 M00004033D:G06 36848 4255 2/24/98 1019 RTA0000043F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA0000043F.a.24.1 M00001455B:A09 24128 4257 3/24/98 5 RTA00000426F.p.04.1 M00004029E:B03 23343 4259 2/24/98 562 RTA00000157.07.1 M00004029C:B03 23343 4260 2/24/98 303					M00004033B:C02	37566
4249 3/24/98 156 RTA00000525F.b.22.1 M00004037C:D07 16679 4250 2/24/98 70 RTA00000409F.m.13.1 M00001618B:E05 0 4251 2/24/98 1198 RTA00000412F.f.10.2 M00003959A:A03 65405 4252 2/24/98 1139 RTA00000404F.h.20.1 M00001619B:A09 15564 4253 3/24/98 41 RTA00000525F.b.17.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000525F.a.22.1 M00004033D:G06 36848 4255 2/24/98 1019 RTA0000043F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA0000043F.g.03.1 M00001455B:A09 24128 4257 3/24/98 5 RTA00000426F.p.04.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000427F.p.07.1 M00004029C:B03 23343 4259 2/24/98 562 RTA00000049F.m.02.1 M00001617A:D06 80933 4261 2/24/98 1201					M00001479D:H03	78382
4250 2/24/98 70 RTA00000409F.m.13.1 M00001618B:E05 0 4251 2/24/98 1198 RTA00000412F.f.10.2 M00003959A:A03 65405 4252 2/24/98 1139 RTA00000404F.h.20.1 M00001619B:A09 15564 4253 3/24/98 41 RTA00000525F.b.17.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000525F.a.22.1 M00004033D:G06 36848 4255 2/24/98 1019 RTA00000403F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA00000403F.a.24.1 M00001479D:G06 23537 4257 3/24/98 5 RTA00000426F.p.04.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029C:B03 23343 4259 2/24/98 562 RTA0000040F.j.17.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA00000409F.m.02.1 M00001616C:A11 9157 4262 3/24/98 241					M00004037C:D07	16679
4251 2/24/98 1198 RTA00000412F.f.10.2 M00003959A:A03 65405 4252 2/24/98 1139 RTA00000404F.h.20.1 M00001619B:A09 15564 4253 3/24/98 41 RTA00000525F.b.17.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000525F.a.22.1 M00004033D:G06 36848 4255 2/24/98 1019 RTA00000403F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA00000426F.p.04.1 M00001455B:A09 24128 4257 3/24/98 5 RTA00000527F.p.07.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029C:B03 23343 4259 2/24/98 562 RTA0000040F.m.02.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA00000527F.o.12.1 M00001616C:A11 9157 4262 3/24/98 241 RTA00000527F.o.12.1 M00001616C:A02 73174 4264 2/24/98 176						0
4252 2/24/98 1139 RTA00000404F.h.20.1 M00001619B:A09 15564 4253 3/24/98 41 RTA00000525F.b.17.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000525F.a.22.1 M00004033D:G06 36848 4255 2/24/98 1019 RTA00000403F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA00000426F.p.04.1 M00001455B:A09 24128 4257 3/24/98 5 RTA00000426F.p.04.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029C:B03 23343 4259 2/24/98 562 RTA00000401F.j.17.1 M00003901B:C05 5483 4260 2/24/98 303 RTA00000130A.h.22.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA000000527F.o.12.1 M00001616C:A11 9157 4262 3/24/98 241 RTA000000527F.o.12.1 M00001616C:A02 73174 4264 2/24/98 176			1198	RTA00000412F.f.10.2	M00003959A:A03	65405
4253 3/24/98 41 RTA00000525F.b.17.1 M00004037B:A04 24715 4254 3/24/98 452 RTA00000525F.a.22.1 M00004033D:G06 36848 4255 2/24/98 1019 RTA00000403F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA00000403F.g.03.1 M00001455B:A09 24128 4257 3/24/98 5 RTA00000426F.p.04.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029C:B03 23343 4259 2/24/98 562 RTA00000527F.p.07.1 M00003901B:C05 5483 4260 2/24/98 303 RTA00000130A.h.22.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA00000409F.m.02.1 M00001616C:A11 9157 4262 3/24/98 241 RTA00000527F.o.12.1 M00001616C:A02 73174 4264 2/24/98 176 RTA0000049F.b.10.1 M00001579D:C03 6539 4265 1/28/98 131		2/24/98		RTA00000404F.h.20.1	M00001619B:A09	15564
4254 3/24/98 452 RTA00000525F.a.22.1 M00004033D:G06 36848 4255 2/24/98 1019 RTA00000403F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA00000403F.a.24.1 M00001455B:A09 24128 4257 3/24/98 5 RTA00000426F.p.04.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029C:B03 23343 4259 2/24/98 562 RTA00000401F.j.17.1 M00003901B:C05 5483 4260 2/24/98 303 RTA00000130A.h.22.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA00000409F.m.02.1 M00001616C:A11 9157 4262 3/24/98 241 RTA00000527F.o.12.1 M00001616C:A02 73174 4264 2/24/98 176 RTA0000049F.b.10.1 M00001455C:G07 73268 4265 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 4266 1/28/98 190					M00004037B:A04	24715
4255 2/24/98 1019 RTA00000403F.g.03.1 M00001479D:G06 23537 4256 2/24/98 532 RTA00000403F.a.24.1 M00001455B:A09 24128 4257 3/24/98 5 RTA00000426F.p.04.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029C:B03 23343 4259 2/24/98 562 RTA00000401F.j.17.1 M00003901B:C05 5483 4260 2/24/98 303 RTA00000130A.h.22.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA00000409F.m.02.1 M00001616C:A11 9157 4262 3/24/98 241 RTA00000527F.o.12.1 M00004028B:G08 688 4263 2/24/98 1170 RTA00000409F.l.24.1 M00001616C:A02 73174 4264 2/24/98 176 RTA00000185AF.d.11.2 M00001579D:C03 6539 4265 1/28/98 131 RTA00000185AR.d.11.1 M00001528A:A01 5175 4266 1/28/98 190			452	RTA00000525F.a.22.1	M00004033D:G06	36848
4256 2/24/98 532 RTA00000403F.a.24.1 M00001455B:A09 24128 4257 3/24/98 5 RTA00000426F.p.04.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029C:B03 23343 4259 2/24/98 562 RTA00000401F.j.17.1 M00003901B:C05 5483 4260 2/24/98 303 RTA00000130A.h.22.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA00000409F.m.02.1 M00001616C:A11 9157 4262 3/24/98 241 RTA00000527F.o.12.1 M00004028B:G08 688 4263 2/24/98 1170 RTA0000049F.l.24.1 M00001616C:A02 73174 4264 2/24/98 176 RTA0000043F.b.10.1 M00001579D:C03 6539 4265 1/28/98 131 RTA00000185AR.d.11.1 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4267 3/24/98 236			1019	RTA00000403F.g.03.1	M00001479D:G06	23537
4257 3/24/98 5 RTA00000426F.p.04.1 M00004029B:H08 34149 4258 3/24/98 43 RTA00000527F.p.07.1 M00004029C:B03 23343 4259 2/24/98 562 RTA00000401F.j.17.1 M00003901B:C05 5483 4260 2/24/98 303 RTA00000130A.h.22.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA00000409F.m.02.1 M00001616C:A11 9157 4262 3/24/98 241 RTA00000527F.o.12.1 M00004028B:G08 688 4263 2/24/98 1170 RTA00000409F.l.24.1 M00001616C:A02 73174 4264 2/24/98 176 RTA00000403F.b.10.1 M00001455C:G07 73268 4265 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236		2/24/98			M00001455B:A09	24128
42583/24/9843RTA00000527F.p.07.1M00004029C:B032334342592/24/98562RTA00000401F.j.17.1M00003901B:C05548342602/24/98303RTA00000130A.h.22.1M00001617A:D068093342612/24/981201RTA00000409F.m.02.1M00001616C:A11915742623/24/98241RTA00000527F.o.12.1M00004028B:G0868842632/24/981170RTA00000409F.l.24.1M00001616C:A027317442642/24/98176RTA00000403F.b.10.1M00001455C:G077326842651/28/98131RTA00000185AF.d.11.2M00001579D:C03653942651/28/98626RTA00000185AR.d.11.1M00001579D:C03653942661/28/98176RTA00000185AR.d.11.1M00001528A:A01517542661/28/98176RTA00000183AF.h.19.1M00001528A:A01517542673/24/9890RTA00000525F.a.03.1M00004031D:F053678642683/24/98236RTA00000527F.o.01.1M00004027A:D0619088					M00004029B:H08	34149
4259 2/24/98 562 RTA00000401F.j.17.1 M00003901B:C05 5483 4260 2/24/98 303 RTA00000130A.h.22.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA00000409F.m.02.1 M00001616C:A11 9157 4262 3/24/98 241 RTA00000527F.o.12.1 M00004028B:G08 688 4263 2/24/98 1170 RTA00000409F.l.24.1 M00001616C:A02 73174 4264 2/24/98 176 RTA00000403F.b.10.1 M00001455C:G07 73268 4265 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 4265 1/28/98 626 RTA00000185AR.d.11.1 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004027A:D06 19088 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088						23343
4260 2/24/98 303 RTA00000130A.h.22.1 M00001617A:D06 80933 4261 2/24/98 1201 RTA00000409F.m.02.1 M00001616C:A11 9157 4262 3/24/98 241 RTA00000527F.o.12.1 M00004028B:G08 688 4263 2/24/98 1170 RTA00000409F.l.24.1 M00001616C:A02 73174 4264 2/24/98 176 RTA00000403F.b.10.1 M00001455C:G07 73268 4265 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 4265 1/28/98 626 RTA00000185AR.d.11.1 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000183AF.h.19.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004027A:D06 19088 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088					M00003901B:C05	5483
4261 2/24/98 1201 RTA00000409F.m.02.1 M00001616C:A11 9157 4262 3/24/98 241 RTA00000527F.o.12.1 M00004028B:G08 688 4263 2/24/98 1170 RTA00000409F.l.24.1 M00001616C:A02 73174 4264 2/24/98 176 RTA00000403F.b.10.1 M00001455C:G07 73268 4265 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 4265 1/28/98 626 RTA00000185AR.d.11.1 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000183AF.h.19.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088				-	M00001617A:D06	
4262 3/24/98 241 RTA00000527F.o.12.1 M00004028B:G08 688 4263 2/24/98 1170 RTA00000409F.l.24.1 M00001616C:A02 73174 4264 2/24/98 176 RTA00000403F.b.10.1 M00001455C:G07 73268 4265 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 4265 1/28/98 626 RTA00000185AR.d.11.1 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000183AF.h.19.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088						
4263 2/24/98 1170 RTA00000409F.l.24.1 M00001616C:A02 73174 4264 2/24/98 176 RTA00000403F.b.10.1 M00001455C:G07 73268 4265 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 4265 1/28/98 626 RTA00000185AR.d.11.1 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000183AF.h.19.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088					M00004028B:G08	688
4264 2/24/98 176 RTA00000403F.b.10.1 M00001455C:G07 73268 4265 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 4265 1/28/98 626 RTA00000185AR.d.11.1 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000183AF.h.19.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088				RTA00000409F.1.24.1	M00001616C:A02	73174
4265 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 4265 1/28/98 626 RTA00000185AR.d.11.1 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000183AF.h.19.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088				RTA00000403F.b.10.1	M00001455C:G07	73268
4265 1/28/98 626 RTA00000185AR.d.11.1 M00001579D:C03 6539 4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000183AF.h.19.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088				RTA00000185AF.d.11.2	M00001579D:C03	6539
4266 1/28/98 190 RTA00000134A.c.7.1 M00001528A:A01 5175 4266 1/28/98 176 RTA00000183AF.h.19.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088						
4266 1/28/98 176 RTA00000183AF.h.19.1 M00001528A:A01 5175 4267 3/24/98 90 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088						
4267 3/24/98 90 RTA00000525F.a.03.1 M00004031D:F05 36786 4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088						
4268 3/24/98 236 RTA00000527F.o.01.1 M00004027A:D06 19088						

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
4270	1/28/98	1 8 3	RTA00000198AF.c.17.1	M00001579C:E08	6923
4271	3/24/98	44	RTA00000527F.p.17.1	M00004030C:D12	17223
4272	3/24/98	129	RTA00000527F.p.18.1	M00004030D:B06	31635
4273	3/24/98	402	RTA00000527F.p.24.1	M00004031B:A06	36832
4274	3/24/98	118	RTA00000525F.a.02.1	M00004031C:H10	37454
4275	1/28/98	353	RTA00000198F.a.9.1	M00001557D:C08	0
4276	2/24/98	1250	RTA00000403F.f.15.1	M00001477D:F10	22768
4277	1/28/98	131	RTA00000185AF.d.11.2	M00001579D:C03	6539
4277	1/28/98	626	RTA00000185AR.d.11.1	M00001579D:C03	6539
4278	2/24/98	428	RTA00000422F.f.14.1	M00001478B:D07	2036
4279	1/28/98	209	RTA00000182AF.c.5.1	M00001464D:F06	6397
4279	1/28/98	304	RTA00000182AR.c.5.1	M00001464D:F06	6397
4280	2/24/98	205	RTA00000138A.n.4.1	M00001624A:G11	21920
4281	2/24/98	251	RTA00000119A.i.9.1	M00001457A:G03	0
4282	3/24/98	250	RTA00000427F.h.24.1	M00004091B:H09	65193
4283	1/28/98	61	RTA00000197AF.h.11.1	M00001476D:G03	22264
4284	3/24/98	216	RTA00000427F.h.11.1	M00004092C:B12	. 26494
4285	1/28/98	110	RTA00000197R.h.01.1	M00001470A:H01	13075
4285	1/28/98	591	RTA00000197AF.h.1.1	M00001470A:H01	13075
4286	1/28/98	110	RTA00000197R.h.01.1	M00001470A:H01	13075
4286	1/28/98	591	RTA00000197AF.h.1.1	M00001470A:H01	13075
4287	3/24/98	276	RTA00000427F.h.19.1	M00004092D:B11	63047
4288	1/28/98	335	RTA00000182AF.e.3.2	M00001468B:H06	0
4289	3/24/98	475	RTA00000427F.i.06.1	M00004097B:D03	41450
4290	2/24/98	286	RTA00000404F.i.19.1	M00001625B:C10	38698
4291	1/28/98	209	RTA00000182AF.c.5.1	M00001464D:F06	6397
4291	1/28/98	304	RTA00000182AR.c.5.1	M00001464D:F06	6397
4292	2/24/98	1165	RTA00000408F.c.08.1	M00001456D:G11	73473
4293	1/28/98	304	RTA00000182AR.c.5.1	M00001464D:F06	6397
4293	1/28/98	209	RTA00000182AF.c.5.1	M00001464D:F06	6397
4294	1/28/98	138	RTA00000182AF.a.3.3	M00001462B:A10	0
4295	1/28/98	36	RTA00000181AF.p.4.3	M00001460A:A03	40392
4296	2/24/98	523	RTA00000418F.j.09.1	M00001626C:D12	76352
4297	2/24/98	296	RTA00000347F.d.06.1	M00001457C:F02	39122
4298	1/28/98	390	RTA00000197AR.f.07.1	M00001457C:C11	19261
4298	1/28/98	184	RTA00000197AF.f.7.1	M00001457C:C11	19261
4299	1/28/98	184	RTA00000197AF.f.7.1	M00001457C:C11	19261
4299	1/28/98	390	RTA00000197AR.f.07.1	M00001457C:C11	19261
4300	1/28/98	133	RTA00000181AR.n.20.3	M00001457B:E03	0
4301	3/24/98	132	RTA00000427F.k.17.1	M00004101A:F07	64965
4302	3/24/98	218	RTA00000427F.i.19.1	M00004102C:D01	64206
4303	3/24/98	436	RTA00000427F.i.21.1	M00004102C:F03	65540
4304	1/28/98	209	RTA00000182AF.c.5.1	M00001464D:F06	6397
4304	1/28/98	304	RTA00000182AR.c.5.1	M00001464D:F06	6397
4305	1/28/98	176	RTA00000183AF.h.19.1	M00001528A:A01	5175
4305	1/28/98	190	RTA00000134A.c.7.1	M00001528A:A01	5175
4306	1/28/98	12	RTA00000183AF.i.15.2	M00001529B:C04	2642
4306	2/24/98	379	RTA00000349R.j.07.1	M00001529B:C04	2642
4307	2/24/98	1156	RTA00000408F.f.10.2	M00001476D:C05	75309
4308	2/24/98	366	RTA00000403F.c.10.1	M00001456D:F05	75261
			227		

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
4309	Appln 2/24/98	Appln	PT 4 00000400F = 17 1	M00001(210 010	7/705
		353 526	RTA00000409F.n.17.1	M00001621C:C10	76725
4310	2/24/98	526	RTA00000411F.a.05.1	M00001675B:H03	76699
4311	2/24/98	90	RTA00000411F.a.02.1	M00001675B:E02	78537
4312	2/24/98	952	RTA00000411F.a.01.1	M00001675B:D02	74524
4313	2/24/98	392	RTA00000410F.p.23.1	M00001675B:C01	73948
4314	2/24/98	238	RTA00000340F.j.12.1	M00001624A:B06	3277
4315	1/28/98	176	RTA00000183AF.h.19.1	M00001528A:A01	5175
4315	1/28/98	190	RTA00000134A.c.7.1	M00001528A:A01	5175
4316	2/24/98	298	RTA00000406F.h.07.1	M00003901B:H04	38003
4317	1/28/98	190	RTA00000134A.c.7.1	M00001528A:A01	5175
4317	1/28/98	176	RTA00000183AF.h.19.1	M00001528A:A01	5175
4318	1/28/98	12	RTA00000183AF.i.15.2	M00001529B:C04	2642
4318	2/24/98	379	RTA00000349R.j.07.1	M00001529B:C04	2642
4319	1/28/98	122	RTA00000197AF.I.15.1	M00001517B:G08	4947
4320	3/24/98	199	RTA00000427F.f.24.1	M00004076D:B09	64572
4321	1/28/98	161	RTA00000183AF.e.23.2	M00001506D:A09	0
4322	1/28/98	17	RTA00000183AR.e.14.2	M00001506B:D09	17437
4323	1/28/98	346	RTA00000197AR.k.22.1	M00001505C:H01	11394
4324	1/28/98	125	RTA00000197AF.k.15.1	M00001504D:D11	22750
4325	1/28/98	212	RTA00000197AF.j.9.1	M00001494B:C01	13236
4326	1/28/98	314	RTA00000182AF.o.5.1	M00001493B:D09	5007
4327	1/28/98	386	RTA00000197AR.j.04.1	M00001492D:A11	17209
4327	1/28/98	259	RTA00000197AF.j.4.1	M00001492D:A11	17209
4328	1/28/98	259	RTA00000197AF.j.4.1	M00001492D:A11	17209
4328	1/28/98	386	RTA00000197AR.j.04.1	M00001492D:A11	17209
4329	1/28/98	94	RTA00000195AF.b.4.1	M00001490C:D07	0
4330	1/28/98	336	RTA00000186AF.f.24.1	M00001629B:E06	0
4330	1/28/98	83	RTA00000186AF.f.24.2	M00001629B:E06	0
4331	2/24/98	1037	RTA00000339F.I.12.1	M00001450A:G11	7711
4332	1/28/98	432	RTA00000198AF.o.05.1	M00003750A:D01	26702
4332	1/28/98	49	RTA00000198R.o.05.1	M00003750A:D01	26702
4333	2/24/98	468	RTA00000423F.c.19.1	M00001680B:E10	40472
4334	2/24/98	1009	RTA00000399F.o.24.1	M00001607D:A11	2272
4335	1/28/98	281	RTA00000188AF.n.10.1	M00003802D:B11	10283
4336 4337	1/28/98	157	RTA00000188AF.n.01.1	M00003801A:B10	36412
4337	2/24/98 2/24/98	842	RTA00000401F.n.23.1	M00003982A:B06	1552
4339	2/24/98	1216	RTA00000404F.e.07.1	M00001608A:D03	9034
4340	2/24/98	1045 4 8 3	RTA00000408F.j.05.2	M00001483C:G06	73878
4341	1/28/98	310	RTA00000406F.g.22.1 RTA00000188AF.m.08.1	M00003881D:C12	38590
4342	1/28/98	118	RTA00000188AF.III.08.1 RTA00000199F.b.24.2	M00003798D:H08	22155
4343	1/28/98	218	RTA00000199F.5.24.2 RTA00000188AF.o.18.1	M00003794A:B03	0
4344	3/24/98	380	RTA00000188AF.0.18.1 RTA00000427F.e.13.1	M00003811D:A12 M00003959D:A04	13678
4345	1/28/98	315	RTA000004271.e.13.1 RTA00000199R.d.23.1		66080
4346	1/28/98	140	RTA00000199K.d.23.1 RTA00000199F.a.2.1	M00003815D:H09 M00003772A:D07	37477
4347	3/24/98	101	RTA00000199F.a.2.1 RTA00000523F.j.19.1		12674
4348	1/28/98	278	RTA00000323F.J.19.1 RTA00000198AF.p.16.1	M00003966B:D02 M00003768A:E02	65910
4349	2/24/98	514	RTA00000198AF.p.10.1 RTA00000404F.e.13.1	M00003768A:E02 M00001608D:E09	71877 12046
4350	1/28/98	508	RTA000004041.e.13.1 RTA00000187AF.i.14.2	M00001608D:E09	12046 19406
4350	2/24/98	928	RTA00000187AT.1.14.2 RTA00000340F.m.04.1	M00001679B:H07	19406
•					17700

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
4351	1/28/98	317	RTA00000198AF.p.09.1	M00003761D:E02	10473
4351	1/28/98	186	RTA00000198R.p.09.1	M00003761D:E02	10473
4352	1/28/98	317	RTA00000198AF.p.09.1	M00003761D:E02	10473
4352	1/28/98	186	RTA00000198R.p.09.1	M00003761D:E02	10473
4353	3/24/98	66	RTA00000427F.b.15.1	M00003971C:F09	66891
4354	1/28/98	508	RTA00000187AF.i.14.2	M00001679B:H07	19406
4354	2/24/98	928	RTA00000340F.m.04.1	M00001679B:H07	19406
4355	1/28/98	144	RTA00000198AF.o.18.1	M00003755A:A09	13018
4356	3/24/98	248	RTA00000527F.l.14.1	M00003983D:A09	14935
4357	1/28/98	347	RTA00000199F.b.03.2	M00003779B:E12	38340
4358	1/28/98	272	RTA00000199F.g.08.2	M00003853D:G08	0
4359	1/28/98	263	RTA00000190AF.n.6.1	M00003965A:B11	0
4360	2/24/98	1183	RTA00000346F.j.21.1	M00003879D:A08	3095
4361	2/24/98	553	RTA00000408F.j.12.2	M00001485B:C03	18226
4362	3/24/98	181	RTA00000523F.b.06.1	M00003808A:F09	28736
4363	1/28/98	246	RTA00000199AF.l.4.1	M00003911D:B04	4410
4364	1/28/98	51	RTA00000199R.k.07.1	M00003901C:A03	12973
4365	1/28/98	62	RTA00000190AF.a.18.2	M00003900D:B10	0
4366	1/28/98	117	RTA00000199AF.j.18.1	M00003889D:B09	5140
4367	1/28/98	255	RTA00000199AF.j.17.1	M00003889A:D10	5121
4368	1/28/98	180	RTA00000199AF.j.12.1	M00003887A:A06	22461
4369	3/24/98	256	RTA00000523F.b.20.1	M00003809C:H07	66492
4370	2/24/98	603	RTA00000399F.o.19.1	M00001607A:F11	2594
4371	2/24/98	510	RTA00000131A.g.16.2	M00001449A:F01	0
4372	1/28/98	49	RTA00000198R.o.05.1	M00003750A:D01	26702
4372	1/28/98	432	RTA00000198AF.o.05.1	M00003750A:D01	26702
4373	2/24/98	424	RTA00000138A.e.13.1	M00001605A:E06	79608
4374	1/28/98	90	RTA00000199F.f.15.2	M00003845A:H12	8772
4375	1/28/98	244	RTA00000199F.f.12.2	M00003844C:A08	8131
4376	1/28/98	78	RTA00000199R.f.09.1	M00003842B:D09	22907
4376	1/28/98	406	RTA00000199F.f.09.2	M00003842B:D09	22907
4377	1/28/98	406	RTA00000199F.f.09.2	M00003842B:D09	22907
4377	1/28/98	78	RTA00000199R.f.09.1	M00003842B:D09	22907
4378	1/28/98	44	RTA00000199F.f.08.2	M00003841D:E03	12445
4379	1/28/98	39	RTA00000189AR.b.19.1	M00003832B:E01	5294
4379	2/24/98	239	RTA00000346F.j.02.1	M00003832B:E01	5294
4380	1/28/98	39	RTA00000189AR.b.19.1	M00003832B:E01	5294
4380	2/24/98	239	RTA00000346F.j.02.1	M00003832B:E01	5294
4381	2/24/98	1161	RTA00000346F.m.05.1	M00003983B:C08	5644
4382	2/24/98	887	RTA00000339F.p.06.1	M00001484A:A10	4880
4383	3/24/98	46	RTA00000523F.c.09.1	M00003813C:D08	47389
4384	2/24/98	1206	RTA00000418F.b.20.1	M00001484D:G05	73560
4385	1/28/98	336	RTA00000186AF.f.24.1	M00001629B:E06	0
4385	1/28/98	83	RTA00000186AF.f.24.2	M00001629B:E06	0
4386	1/28/98	111	RTA00000198AF.o.12.1	M00003751D:B02	22038 1015
4387	3/24/98	365	RTA00000527F.k.16.1	M00003982B:B06	
4388	2/24/98	1113	RTA00000418F.p.21.1	M00001677D:F03 M00003982B:H07	78068 17148
4389	3/24/98	281	RTA00000527F.k.20.1 RTA00000198F.i.5.1	M00003982B:H07 M00001638A:D10	17148 39989
4390	1/28/98	360 55	RTA00000198F.I.5.1	M00001638A:D10 M00001636C:H09	6033
4391	1/28/98	33	KIAUUUUI 60AF.I.21.1	14100001036C:HU9	0033

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appln			
4392	1/28/98	316	RTA00000198AF.h.24.1	M00001636C:C01	8390
4393	1/28/98	208	RTA00000198AF.h.22.1	M00001635C:A03	22366
4394	2/24/98	1031	RTA00000404F.g.08.1	M00001613D:H10	38980
4395	3/24/98	382	RTA00000427F.a.12.1	M00003982C:H10	63377
4396	2/24/98	916	RTA00000418F.p.20.1	M00001677D:B07	78023
4397	1/28/98	91	RTA00000198AF.j.19.1	M00001653C:F12	38914
4398	2/24/98	858	RTA00000341F.e.20.1	M00003891D:B10	67422
4399	1/28/98	354	RTA00000198R.k.03.1	M00001655A:F06	22765
4399	1/28/98	158	RTA00000198AF.k.03.1	M00001655A:F06	22765
4400	3/24/98	219	RTA00000427F.f.17.1	M00004115A:B12	63803
4401	3/24/98	153	RTA00000527F.I.13.1	M00003983C:F10	36904
4402	3/24/98	320	RTA00000427F.j.06.1	M00004102D:B05	63676
4403	2/24/98	762	RTA00000411F.c.04.1	M00001677B:E06	76858
4404	2/24/98	957	RTA00000411F.c.03.1	M00001677B:B06	79280
4405	3/24/98	479	RTA00000527F.I.23.1	M00003984A:B06	36018
4406	1/28/98	329	RTA00000186AF.b.9.1	M00001616C:F07	0
4407	2/24/98	1115	RTA00000340F.i.08.1	M00001615B:F07	12005
4408	2/24/98	1022	RTA00000401F.j.15.1	M00003901A:C09	3061
4409	1/28/98	4	RTA00000198R.f.04.1	M00001607D:F07	5023
4410	3/24/98	173	RTA00000426F.m.18.1	M00003986D:G07	62974
4411	2/24/98	616	RTA00000423F.c.11.1	M00001677D:B02	0
4412	1/28/98	345	RTA00000187AF.h.21.1	M00001679A:F01	39171
4413	2/24/98	621	RTA00000408F.i.18.2	M00001482C:D02	74410
4414	1/28/98	344	RTA00000198AF.o.02.1	M00003748A:B07	68756
4415	2/24/98	257	RTA00000411F.c.17.1	M00001678D:G03	77664
4416	2/24/98	944	RTA00000422F.k.24.1	M00001610C:E06	39118
4417	2/24/98	876	RTA00000423F.d.16.1	M00001678D:C11	39173
4418	2/24/98	1144	RTA00000345F.j.09.1	M00001451B:F01	13
4419	1/28/98	242	RTA00000198AF.m.17.1	M00001679D:F06	77992
4419	1/28/98	260	RTA00000198R.m.17.1	M00001679D:F06	77992
4420	1/28/98	260	RTA00000198R.m.17.1	M00001679D:F06	77992
4420	1/28/98	242	RTA00000198AF.m.17.1	M00001679D:F06	77992
4421	1/28/98	242	RTA00000198AF.m.17.1	M00001679D:F06	77992
4421	1/28/98	260	RTA00000198R.m.17.1	M00001679D:F06	77992
4422	1/28/98	260	RTA00000198R.m.17.1	M00001679D:F06	77992
4422	1/28/98	242	RTA00000198AF.m.17.1	M00001679D:F06	77992
4423	1/28/98	238	RTA00000187AR.k.12.1	M00001679D:F02	78415
4423	2/24/98	407	RTA00000340R.m.07.1	M00001679D:F02	78415
4424	1/28/98	276	RTA00000198AF.j.15.1	M00001653B:E09	4369
4425	1/28/98	65	RTA00000198AF.m.16.1	M00001679D:D05	51
4426	1/28/98	257	RTA00000198AF.e.20.1	M00001604C:E09	9810
4427	2/24/98	820	RTA00000423F.d.11.1	M00001678C:C06	38950
4428	3/24/98	466	RTA00000427F.d.06.1	M00003980B:C06	33446
4429	2/24/98	455	RTA00000399F.d.23.1	M00001481B:A07	3310
4430	2/24/98	851	RTA00000423F.d.07.1	M00001678B:B12	0
4431	1/28/98	142	RTA00000198AF.k.19.1	M00001660B:C04	75879
4432	2/24/98	485	RTA00000419F.a.02.1	M00001678A:F05	77993
4433	3/24/98	224	RTA00000527F.k.09.1	M00003981C:F05	213
4434	2/24/98	1032	RTA00000423F.c.13.1	M00001678A:A11	39059
4435	1/28/98	354	RTA00000198R.k.03.1	M00001655A:F06	22765

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln		14000014551 F04	22765
4435	1/28/98	158	RTA00000198AF.k.03.1	M00001655A:F06	22765
4436	1/28/98	354	RTA00000198R.k.03.1	M00001655A:F06	22765
4436	1/28/98	158	RTA00000198AF.k.03.1	M00001655A:F06	22765
4437	1/28/98	158	RTA00000198AF.k.03.1	M00001655A:F06	22765
4437	1/28/98	354	RTA00000198R.k.03.1	M00001655A:F06	22765
4438	1/28/98	238	RTA00000187AR.k.12.1	M00001679D:F02	78415
4438	2/24/98	407	RTA00000340R.m.07.1	M00001679D:F02	78415
4439	1/28/98	669	RTA00000192AF.c.2.1	M00004121B:G01	0
4440	3/24/98	394	RTA00000527F.g.14.1	M00003845D:B02	37532
4441	1/28/98	608	RTA00000192AF.p.8.1	M00004212B:C07	2379
4441	2/24/98	653	RTA00000352R.m.12.1	M00004212B:C07	2379
4442	1/28/98	608	RTA00000192AF.p.8.1	M00004212B:C07	2379
4442	2/24/98	653	RTA00000352R.m.12.1	M00004212B:C07	2379
4443	1/28/98	730	RTA00000192AF.o.11.1	M00004205D:F06	0
4444	2/24/98	1157	RTA00000422F.m.18.1	M00001647B:E04	23829
4445	2/24/98	1187	RTA00000120A.c.19.1	M00001464A:B03	81016
4446	2/24/98	913	RTA00000120A.c.20.1	M00001464A:B07	43235
4447	1/28/98	589	RTA00000192AF.l.1.1	M00004183C:D07	16392
4448	2/24/98	640	RTA00000405F.f.02.1	M00001669B:G02	38665
4446 4449	1/28/98	27	RTA00000192AF.i.12.1	M00004169C:C12	5319
	2/24/98	6 8 1	RTA0000013211211	M00001464A:D03	34278
4450	2/24/98	265	RTA0000012071.6.2411	M00001647B:C09	13157
4451	1/28/98	70	RTA00000192AF.e.3.1	M00004138B:H02	13272
4452	3/24/98	70 171	RTA00000172A1.c.3.1	M00003829A:F03	62878
4453	3/24/98 2/24/98	1134	RTA000003231.c.10.1	M00001650A:A12	74550
4454		618	RTA00000192AF.a.14.1	M00004111D:A08	6874
4455	1/28/98 1/28/98	457	RTA00000192AI.a.14.1 RTA00000191AR.1.7.2	M00004081C:D12	14391
4456		596	RTA00000191AR.i.7.2	M00003846B:D06	6874
4457	2/24/98 3/24/98	460	RTA00000537K.f.05.1	M00003840B:E07	26522
4458		400	RTA00000523F.f.17.1	M00003840B:E08	63984
4459	3/24/98	400 1129	RTA000003231.1.17.1 RTA00000401F.m.07.1	M00003917D:E11	2893
4460	2/24/98	132	RTA000004117.m.07.1	M00001650B:C10	73600
4461	2/24/98	482	RTA00000187AF.j.7.1	M00001679C:F01	78091
4462	1/28/98	4 6 2 1107	RTA00000187A1 3.7.1	M00003903D:C06	78444
4463	2/24/98	609	RTA000004191.i.22.1 RTA00000404F.o.10.2	M00003563B:B12	16785
4464	2/24/98	376	RTA00000177AF.m.18.3	M00001355B:G11	0
4465	1/28/98		RTA00000177AF.m.18.1	M00001355B:G11	0
4465	1/28/98	375	RTA00000177A1.iii.18.1	M00001353B:G17	81284
4466	2/24/98	186	RTA00000192A.k.o.1 RTA00000196AF.c.17.1	M00001352C:F06	39602
4467	1/28/98	18	RTA00000130A1.c.17.1	M00004108C:E01	64547
4468	3/24/98	282 859	RTA000004271.ii.22.1 RTA00000419F.m.22.1	M00003914A:G09	75600
4469	2/24/98	33	RTA000004191.iii.22.ii RTA00000524F.b.21.1	M0000531171:309	0
4470	3/24/98		RTA00000524F.b.21.1 RTA00000523F.d.12.1	M000032103:D08	64888
4471	3/24/98	170	RTA00000523F.d.12.1 RTA00000523F.d.18.1	M00003822B:G01	64072
4472	3/24/98	117	RTA00000323F.d.18.1 RTA00000423F.h.20.1	M00003914A:G06	38639
4473	2/24/98	739 527	RTA00000423F.ii.20.1 RTA00000419F.m.21.1	M00003914A:E04	77947
4474	2/24/98	527	RTA00000419F.iii.21.1 RTA00000119A.j.22.1	M0000371471:E04 M00001460A:F07	80336
4475	2/24/98	237	RTA00000119A.J.22.1 RTA00000404F.m.10.2	M000014007:107 M00001641D:E02	779
4476	2/24/98	349 462	RTA00000404F.iii.10.2 RTA00000119A.j.23.1	M00001041D:E02	79835
4477	2/24/98	462	RTA00000119A.J.23.1 RTA00000341F.i.22.1	M0000140071:G07	7825
4478	2/24/98	1263	NIA000003411.1.22.1	1410000007111111111	. 020

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
110.	Priority	Priority			
	Appln	Appln			
4479	3/24/98	47	RTA00000523F.e.18.1	M00003829D:A11	62898
4480	1/28/98	152	RTA00000196AF.c.20.1	M00001352C:H02	8934
4481	3/24/98	13	RTA00000528F.m.16.1	M00003845D:C03	4468
4482	1/28/98	14	RTA00000196R.c.11.2	M00001352A:E12	13658
4483	2/24/98	641	RTA00000410F.j.20.1	M00001642D:G10	73601
4484	1/28/98	141	RTA00000196AF.c.6.1	M00001350A:D06	23148
4485	1/28/98	25	RTA00000196AF.c.1.1	M00001349C:C05	8171
4486	2/24/98	436	RTA00000119A.m.15.1	M00001461A:E05	80989
4487	1/28/98	9	RTA00000177AF.g.22.1	M00001347C:G08	7031
4488	2/24/98	162	RTA00000406F.1.08.1	M00003908D:D12	39016
4489	2/24/98	1056	RTA00000419F.m.18.1	M00003908C:G09	76014
4490	1/28/98	73	RTA00000177AF.e.21.3	M00001344A:H07	4306
4491	3/24/98	326	RTA00000527F.e.09.1	M00003826B:E11	37521
4492	2/24/98	900	RTA00000419F.m.13.1	M00003908A:F12	79052
4493	2/24/98	441	RTA00000404F.m.20.2	M00001647A:H08	39144
4494	2/24/98	1217	RTA00000410F.j.17.1	M00001642D:F02	72912
4495	3/24/98	309	RTA00000523F.j.10.1	M00003860B:G09	63384
4496	2/24/98	385	RTA00000418F.m.14.1	M00001651B:E06	75711
4497	2/24/98	1121	RTA00000120A.m.10.3	M00001467A:B03	81376
4498	1/28/98	617	RTA00000179AF.d.13.3	M00001394A:F01	6583
4499	2/24/98	1242	RTA00000405F.d.10.1	M00001661C:F11	39000
4500	3/24/98	19	RTA00000527F.j.02.2	M00003856A:B07	4896
4501	2/24/98	645	RTA00000422F.p.12.2	M00001661C:F10	9840
4502	3/24/98	142	RTA00000523F.i.18.1	M00003856B:C04	64463
4503	2/24/98	376	RTA00000400F.k.22.1	M00001656A:B07	2512
4504	1/28/98	532	RTA00000177AF.o.4.1	M00001358C:C06	0
4505	2/24/98	1128	RTA00000423F.a.02.3	M00001656B:A08	39210
4506	2/24/98	1143	RTA00000423F.a.03.1	M00001656B:D05	26796
4507	2/24/98	408	RTA00000405F.d.14.1	M00001662A:C12	35209
4508	3/24/98	360	RTA00000523F.j.03.1	M00003860A:A08	64535
4509	1/28/98	409	RTA00000180AF.d.1.3	M00001418D:B06	8526
4510	2/24/98	784	RTA00000418F.o.14.1	M00001661B:B05	33524
4511	3/24/98	120	RTA00000426F.h.09.1	M00003905B:G03	78797
4512	1/28/98	706	RTA00000177AF.i.6.4	M00001350A:B08	0
4513	3/24/98	4	RTA00000426F.h.11.1	M00003905B:H05	75479
4514	2/24/98	697	RTA00000412F.d.14.1	M00003905D:C08	76757
4515	2/24/98	908	RTA00000423F.g.03.1	M00003905C:G11	38007
4516	3/24/98	342	RTA00000427F.e.12.1	M00003959C:G06	62813
4517	2/24/98	97	RTA00000403F.e.01.1	M00001473A:C11	38965
4518	2/24/98	555	RTA00000133A.d.22.1	M00001469A:G11	11797
4519	2/24/98	454	RTA00000418F.n.19.1	M00001659C:F02	28761
4520	2/24/98	562	RTA00000401F.j.17.1	M00003901B:C05	5483
4521	2/24/98	1215	RTA00000422F.o.08.2	M00001659D:D03	26832
4522	2/24/98	635	RTA00000418F.o.17.1	M00001661B:F03	79069
4523	3/24/98	190	RTA00000523F.h.12.1	M00003851C:D07	65745
4524	1/28/98	267	RTA00000186AF.g.11.2	M00001630B:H09	5214
4525	2/24/98	238	RTA00000340F.j.12.1	M00001624A:B06	3277
4526	2/24/98	331	RTA00000404F.o.18.2	M00001651C:C05	39110
4527	1/28/98	626	RTA00000185AR.d.11.1	M00001579D:C03	6539
4527	1/28/98	131	RTA00000185AF.d.11.2	M00001579D:C03	6539

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
4500	Appln 1/28/98	Appln 626	RTA00000185AR.d.11.1	M00001579D:C03	6539
4528 4528	1/28/98	131	RTA00000185AF.d.11.2	M00001579D:C03	6539
	1/28/98	147	RTA00000185AF.c.24.2	M00001578B:E04	23001
4529	2/24/98	164	RTA00000185A1.c.24.2	M00001370B:201	0
4530		611	RTA000003431 .k.00.1 RTA00000404F.p.02.2	M0000147371.712	39097
4531	2/24/98	274	RTA00000404F.p.02.2 RTA00000405F.e.08.1	M00001652D:7100	37916
4532	2/24/98		RTA00000403F.e.08.1 RTA00000423F.d.17.1	M00001663A:C11	20630
4533	2/24/98	755	RTA00000423F.d.17.1 RTA00000422F.j.20.1	M000016653A:G07	22388
4534	2/24/98	126	RTA00000422F.j.20.1 RTA00000523F.i.08.1	M00001055A:G07	65099
4535	3/24/98	195	RTA00000527F.i.05.1	M00003853A:C12	37481
4536	3/24/98	83	RTA00000327F.1.03.2 RTA00000177AF.m.18.1	M00003851C.B00	0
4537	1/28/98	375		M00001355B:G11	0
4537	1/28/98	376	RTA00000177AF.m.18.3	M00001535B:G11 M00001545A:C03	19255
4538	2/24/98	763	RTA00000135A.m.18.1	M00001543A:C03	74986
4539	2/24/98	362	RTA00000418F.m.16.1	M00001653B.E00 M00001662C:A04	11736
4540	2/24/98	287	RTA00000410F.n.09.1		0
4541	3/24/98	416	RTA00000527F.i.12.2	M00003852B:D11	=
4542	2/24/98	662	RTA00000339F.o.07.1	M00001473D:G01	2566
4543	2/24/98	949	RTA00000340R.j.07.1	M00001654C:D05	38954
4544	3/24/98	146	RTA00000527F.i.17.2	M00003853B:C08	37539
4545	2/24/98	939	RTA00000405F.a.03.1	M00001654C:E04	39065
4546	3/24/98	42	RTA00000527F.i.19.2	M00003853C:C06	38089
4547	3/24/98	381	RTA00000426F.f.18.1	M00003854C:C02	63271
4548	2/24/98	656	RTA00000403F.e.08.1	M00001473D:B11	19126
4549	3/24/98	37	RTA00000426F.f.20.1	M00003854C:F01	65134
4550	2/24/98	733	RTA00000405F.d.18.1	M00001662C:B02	10494
4551	1/28/98	334	RTA00000181AR.b.21.3	M00001444C:D05	3266
4551	1/28/98	321	RTA00000181AR.b.21.1	M00001444C:D05	3266
4552	3/24/98	198	RTA00000523F.o.05.1	M00005175B:H04	0
4553	3/24/98	302	RTA00000427F.p.04.2	M00005100B:H07	0
4554	3/24/98	203	RTA00000427F.p.10.2	M00005102C:F09	0
4555	1/28/98	331	RTA00000197AR.c.20.1	M00001449D:A06	16282
4556	3/24/98	6	RTA00000523F.l.10.1	M00005134B:E01	0
4557	1/28/98	174	RTA00000181AF.e.22.3	M00001448D:F09	3442
4558	3/24/98	79	RTA00000523F.I.15.1	M00005134C:E11	0
4559	3/24/98	386	RTA00000523F.I.16.1	M00005134C:G04	0
4560	3/24/98	76	RTA00000523F.l.18.1	M00005134D:A06	0
4561	3/24/98	192	RTA00000523F.m.02.1	M00005134D:H03	0
4562	3/24/98	290	RTA00000427F.1.03.1	M00005136D:B07	0
4563	3/24/98	269	RTA00000427F.p.02.2	M00005100B:D02	0
4564	1/28/98	321	RTA00000181AR.b.21.1	M00001444C:D05	3266
4564	1/28/98	334	RTA00000181AR.b.21.3	M00001444C:D05	3266
4565	3/24/98	334	RTA00000427F.n.11.1	M00004960B:A09	0
4566	1/28/98	334	RTA00000181AR.b.21.3	M00001444C:D05	3266
4566	1/28/98	321	RTA00000181AR.b.21.1	M00001444C:D05	3266
4567	1/28/98	334	RTA00000181AR.b.21.3	M00001444C:D05	3266
4567	1/28/98	321	RTA00000181AR.b.21.1	M00001444C:D05	3266
4568	3/24/98	328	RTA00000523F.n.01.1	M00005137A:E01	0
4569	1/28/98	356	RTA00000180AF.I.12.2	M00001433B:H11	0
4570	3/24/98	68	RTA00000523F.n.04.1	M00005138B:D12	0
4571	3/24/98	127	RTA00000523F.n.10.1	M00005140D:G09	0

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appin	Appln	DT 4 00000100 4 D : 04 4	M00001429C:G12	22300
4572	1/28/98	187	RTA00000180AR.j.04.4	M00001429C:G12 M00005173C:A02	0
4573	3/24/98	112	RTA00000523F.n.12.1		0
4574	3/24/98	305	RTA00000523F.n.16.1	M00005173D:H02 M00005174D:B02	0
4575	3/24/98	164	RTA00000523F.n.17.1	M00005174D:B02 M00005174D:H02	0
4576	3/24/98	107	RTA00000523F.n.20.1	M00003174D.H02 M00001641C:C06	79058
4577	2/24/98	898	RTA00000418F.I.03.1	M00001641C:C06 M00005136D:C01	0
4578	3/24/98	288	RTA00000427F.1.04.1		0
4579	3/24/98	462	RTA00000427F.p.13.2	M00004695B:E04	13238
4580	1/28/98	137	RTA00000181AF.m.4.3	M00001455A:E09	2364
4581	1/28/98	20	RTA00000181AF.l.14.2	M00001454D:D06	
4582	3/24/98	105	RTA00000526F.d.01.1	M00004104B:A02	4468
4583	3/24/98	261	RTA00000427F.i.22.1	M00004104D:B05	63199
4584	3/24/98	81	RTA00000427F.j.07.1	M00004105A:B10	64819
458 5	3/24/98	287	RTA00000525F.d.19.1	M00004114B:D09	36860
4586	1/28/98	311	RTA00000191AR.j.4.2	M00004071D:A10	5198
4587	3/24/98	337	RTA00000525F.e.08.1	M00004115C:H04	24193
4588	3/24/98	206	RTA00000525F.f.07.1	M00004119A:A06	37500
4589	3/24/98	461	RTA00000427F.f.15.1	M00004119D:A07	66734
4590	3/24/98	410	RTA00000427F.f.16.1	M00004119D:H06	64122
4591	3/24/98	307	RTA00000427F.p.03.2	M00005100B:G11	0
4592	3/24/98	180	RTA00000523F.k.02.1	M00004687A:C03	0
4593	1/28/98	115	RTA00000179AR.o.20.3	M00001409D:F11	2409
4594	3/24/98	315	RTA00000427F.n.19.1	M00004891D:E07	0
4595	3/24/98	375	RTA00000427F.p.19.2	M00004895C:G05	0
4596	3/24/98	470	RTA00000427F.p.24.2	M00004897D:F03	0
4597	1/28/98	155	RTA00000197F.e.8.1	M00001454A:C11	3135
4598	1/28/98	286	RTA00000181AR.k.2.3	M00001453C:A11	0
4598	1/28/98	389	RTA00000181AR.k.2.2	M00001453C:A11	0
4599	1/28/98	286	RTA00000181AR.k.2.3	M00001453C:A11	0
4599	1/28/98	389	RTA00000181AR.k.2.2	M00001453C:A11	0
4600	1/28/98	285	RTA00000181AR.j.14.3	M00001453B:E10	5399
4601	3/24/98	317	RTA00000428F.a.01.1	M00004897D:G05	0
4602	3/24/98	85	RTA00000427F.m.21.1	M00004900C:E11	0
4603	3/24/98	121	RTA00000427F.n.02.1	M00004900D:B10	0
4604	3/24/98	78	RTA00000427F.o.05.1	M00004958B:D01	0
4605	3/24/98	437	RTA00000427F.n.10.1	M00004960B:A08	0
4606	3/24/98	388	RTA00000526F.d.17.1	M00004235A:A12	2757
4607	1/28/98	299	RTA00000196AF.f.5.1	M00001366D:G02	11937
4608	1/28/98	369	RTA00000196F.m.3.1	M00001413A:F02	10453
4609	3/24/98	319	RTA00000523F.p.15.1	M00005178B:H01	0
4610	1/28/98	374	RTA00000178AF.I.11.1	M00001383A:G04	23286
4611	2/24/98	1090	RTA00000405F.g.22.1	M00001673C:A02	527
4612	1/28/98	127	RTA00000178AF.k.18.1	M00001382A:F04	9755
4613	1/28/98	104	RTA00000196R.h.03.1	M00001381A:D02	6636
4614	2/24/98	642	RTA00000341F.h.19.1	M00003916C:C05	0
4615	2/24/98	655	RTA00000351R.p.14.1	M00003915C:H04	13166
4616	1/28/98	145	RTA00000178AF.h.24.1	M00001376B:C06	6745
4617	2/24/98	224	RTA00000341F.g.21.1	M00003914C:F09	8823
4618	2/24/98	301	RTA00000401F.m.23.1	M00003914C:C02	2801
4619	2/24/98	133	RTA00000404F.1.20.1	M00001639B:H05	38638

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln	DC 4 00000404E 1 20 2	M00001620D-U05	38638
4619	2/24/98	63	RTA00000404F.I.20.2	M00001639B:H05 M00001641B:C10	78988
4620	2/24/98	542	RTA00000410F.i.19.1	M00001641B:C10 M00001639B:H05	38638
4621	2/24/98	63	RTA00000404F.1.20.2	M00001639B:H05	38638
4621	2/24/98	133	RTA00000404F.1.20.1		14959
4622	2/24/98	600	RTA00000406F.m.04.1	M00003914B:A11	945
4623	1/28/98	33	RTA00000178AR.a.20.1	M00001362C:H11	943 945
4623	2/24/98	979	RTA00000345F.b.17.1	M00001362C:H11	
4624	1/28/98	33	RTA00000178AR.a.20.1	M00001362C:H11	945
4624	2/24/98	979	RTA00000345F.b.17.1	M00001362C:H11	945
4625	3/24/98	73	RTA00000524F.b.12.1	M00005213C:G01	0
4626	1/28/98	373	RTA00000196F.e.12.1	M00001361C:H11	10147
4627	2/24/98	1233	RTA00000418F.l.02.1	M00001641C:C05	39316
4628	3/24/98	184	RTA00000524F.b.18.1	M00005214B:D11	0
4629	3/24/98	353	RTA00000428F.a.18.1	M00005214C:A09	0
4630	1/28/98	89	RTA00000177AF.n.8.3	M00001356D:F06	4188
4630	1/28/98	15	RTA00000177AR.n.8.1	M00001356D:F06	4188
4631	1/28/98	89	RTA00000177AF.n.8.3	M00001356D:F06	4188
4631	1/28/98	15	RTA00000177AR.n.8.1	M00001356D:F06	4188
4632	1/28/98	375	RTA00000177AF.m.18.1	M00001355B:G11	0
4632	1/28/98	376	RTA00000177AF.m.18.3	M00001355B:G11	0
4633	1/28/98	375	RTA00000177AF.m.18.1	M00001355B:G11	0
4633	1/28/98	376	RTA00000177AF.m.18.3	M00001355B:G11	0
4634	2/24/98	682	RTA00000410F.i.17.1	M00001641B:B01	78147
4635	1/28/98	367	RTA00000196F.i.24.1	M00001392C:D10	4233
4636	1/28/98	264	RTA00000179AF.k.3.3	M00001401A:H07	0
4637	1/28/98	333	RTA00000196F.k.15.1	M00001400A:F06	8320
4638	1/28/98	38	RTA00000196R.k.07.1	M00001399C:D09	22443
4638	1/28/98	289	RTA00000196F.k.07.1	M00001399C:D09	22443
4639	1/28/98	38	RTA00000196R.k.07.1	M00001399C:D09	22443
4639	1/28/98	289	RTA00000196F.k.07.1	M00001399C:D09	22443
4640	1/28/98	289	RTA00000196F.k.07.1	M00001399C:D09	22443
4640	1/28/98	38	RTA00000196R.k.07.1	M00001399C:D09	22443
4641	1/28/98	38	RTA00000196R.k.07.1	M00001399C:D09	22443
4641	1/28/98	289	RTA00000196F.k.07.1	M00001399C:D09	22443
4642	3/24/98	324	RTA00000523F.o.09.1	M00005176A:C12	0
4643	3/24/98	122	RTA00000523F.o.12.1	M00005177A:B06	0
4644	1/28/98	167	RTA00000179AF.d.22.3	M00001394C:C11	7955
4645	1/28/98	351	RTA00000179AF.c.22.1	M00001393B:B09	22515
4645	1/28/98	459	RTA00000179AF.c.22.3	M00001393B:B09	22515
4646	1/28/98	351	RTA00000179AF.c.22.1	M00001393B:B09	22515
4646	1/28/98	459	RTA00000179AF.c.22.3	M00001393B:B09	22515
4647	3/24/98	361	RTA00000523F.p.08.1	M00005178A:A07	0
4648	1/28/98	43	RTA00000179AF.c.14.3	M00001392D:H04	0
4649	3/24/98	268	RTA00000177711.6.11.15	M00004103B:B07	62851
4649 4650	3/24/98	473	RTA000004271.k.19.1 RTA00000523F.o.21.1	M00005177C:A01	0
4650 4651	1/28/98	60	RTA000003251.0.21.1 RTA00000196AR.i.12.3	M00001389D:G11	38800
4651 4651	1/28/98	128	RTA00000196R.i.12.1	M00001389D:G11	38800
	1/28/98	128	RTA00000196F.i.12.1	M00001389D:G11	38800
4652	1/28/98	60	RTA000001961.1.12.1 RTA00000196AR.i.12.3	M00001389D:G11	38800
4652	1/28/98	60	RTA00000196AR.i.12.3	M00001389D:G11	38800
4653	1/20/98	OU	K 1 A00000 1 70AK.1. 12.3		

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
NO:	Priority	Priority			
	-	Appln			
4653	Appln 1/28/98	128	RTA00000196F.i.12.1	M00001389D:G11	38800
4654	1/28/98	60	RTA00000196AR.i.12.3	M00001389D:G11	38800
4654	1/28/98	128	RTA00000196F.i.12.1	M00001389D:G11	38800
4655	1/28/98	28	RTA000001781.12.11	M00001387B:H07	0
	1/28/98	279	RTA00000176/IR.0.01.5	M00001386A:D11	7308
4656	1/28/98	130	RTA00000196AF.h.23.1	M00001386A:C02	13357
4657	1/28/98	254	RTA00000178AF.n.2.1	M00001385C:H11	17083
4658	1/28/98	234 74	RTA00000176AF.h.20.1	M00001385B:F10	0
4659		377	RTA00000170AF.m.19.1	M00001384D:H07	0
4660	1/28/98	120	RTA00000178AR.m.19.5	M00001384D:H07	Ö
4660	1/28/98	377	RTA00000178AF.m.19.1	M00001384D:H07	ő
4661	1/28/98	120	RTA00000178AR.m.19.5	M00001384D:H07	ŏ
4661	1/28/98	228	RTA00000178AR.iii.17.3	M00005177A:H09	Ö
4662	3/24/98		RTA000003231.0.14.1 RTA00000177AR.m.13.1	M00003177A:1103 M00001355A:C12	4175
4663	1/28/98	567	RTA00000177AR.m.13.1 RTA00000177AR.m.13.3	M00001355A:C12	4175
4663	1/28/98	538	RTA00000177AR.m.13.3 RTA00000177AR.m.13.4	M00001355A:C12	4175
4663	1/28/98	533		M00001333A:C12	12498
4664	1/28/98	511	RTA00000196AF.g.10.1	M00001370B:A02 M00004692A:E07	0
4665	1/28/98	620	RTA00000201R.g.08.1	M00004692A:E07	0
4665	1/28/98	619	RTA00000201F.g.08.1	M00004692A:E07	0
4665	1/28/98	621	RTA00000201R.g.08.2	M00004652A.E07	74341
4666	3/24/98	194	RTA00000522F.j.12.2	M00001831C:A04 M00003842C:D11	66700
4667	2/24/98	79	RTA00000419F.g.08.1	M00003842C.D11 M00004692A:E07	00700
4668	1/28/98	619	RTA00000201F.g.08.1	M00004692A:E07	0
4668	1/28/98	620	RTA00000201R.g.08.1	M00004692A:E07	0
4668	1/28/98	621	RTA00000201R.g.08.2	M00004092A.E07	3114
4669	1/28/98	529	RTA00000178AF.b.13.1	M00001560A:F03	9900
4670	2/24/98	111	RTA00000128A.i.20.1	M00001300A.F03	77622
4671	3/24/98	379	RTA00000522F.k.02.2	M00001652D:B09	77619
4672	3/24/98	135	RTA00000522F.k.10.2	M00001550A:H06	80085
4673	2/24/98	1197	RTA00000128A.j.10.1	M00001560A:H10	5316
4674	2/24/98	140	RTA00000128A.j.6.2	M00001300A.1110	77373
4675	3/24/98	247	RTA00000425F.j.21.1 RTA00000177AR.m.13.3	M00001055B:B11	4175
4676	1/28/98	538		M00001355A:C12	4175
4676	1/28/98	567	RTA00000177AR.m.13.1 RTA00000177AR.m.13.4	M00001355A:C12	4175
4676	1/28/98	533	RTA00000177AR.ii.13.4 RTA00000403F.m.20.1	M00001576A:F11	707
4677	2/24/98	729 427	RTA00000403F.m.20.1	M00001576A:F11	707
4677	2/24/98	437	RTA00000403F.III.20.2 RTA00000177AR.m.13.4	M00001376A:111	4175
4678	1/28/98	533	RTA00000177AR.m.13.4	M00001355A:C12	4175
4678	1/28/98	538	RTA00000177AR.m.13.1	M00001355A:C12	4175
4678	1/28/98	567	RTA00000177AR.m.13.4	M00001355A:C12	4175
4679	1/28/98	533	RTA00000177AR.m.13.4 RTA00000177AR.m.13.3	M00001355A:C12	4175
4679	1/28/98	538	RTA00000177AR.m.13.3	M00001355A:C12	4175
4679	1/28/98	567	RTA00000177AR.m.13.1 RTA00000177AR.m.13.3	M00001355A:C12	4175
4680	1/28/98	538 567	RTA00000177AR.m.13.3	M00001355A:C12	4175
4680	1/28/98			M00001355A:C12	4175
4680	1/28/98	533	RTA00000177AR.m.13.4 RTA00000177AR.m.13.4	M00001355A:C12	4175
4681	1/28/98	533	RTA00000177AR.m.13.4 RTA00000177AR.m.13.3	M00001355A:C12	4175
4681	1/28/98	538	RTA00000177AR.m.13.3 RTA00000177AR.m.13.1	M00001355A:C12	4175
4681	1/28/98	567 620	RTA000001//AR.m.13.1 RTA00000201R.g.08.1	M00001333A.C12 M00004692A:E07	0
4682	1/28/98	620	-	1410000T072/A.LU/	J
			236		

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority Appln			
4682	Appln 1/28/98	621	RTA00000201R.g.08.2	M00004692A:E07	0
4682	1/28/98	619	RTA00000201F.g.08.1	M00004692A:E07	0
4683	2/24/98	845	RTA00000348R.b.04.1	M00001342B:E01	1890
4684	2/24/98	1088	RTA00000339R.b.02.1	M00001344B:F12	0
4684	2/24/98	1149	RTA00000339F.b.02.1	M00001344B:F12	0
4685	2/24/98	618	RTA00000423F.1.04.1	M00004039B:G08	14320
4686	2/24/98	1235	RTA00000411F.j.04.1	M00003841C:F03	66219
4687	2/24/98	415	RTA00000340R.f.05.1	M00001569B:G11	3202
4688	2/24/98	1088	RTA00000339R.b.02.1	M00001344B:F12	0
4688	2/24/98	1149	RTA00000339F.b.02.1	M00001344B:F12	0
4689	2/24/98	1067	RTA00000411F.j.11.1	M00003841D:F06	66154
4690	1/28/98	429	RTA00000196F.i.19.1	M00001390C:C11	39498
4690	2/24/98	925	RTA00000353R.h.10.1	M00001390C:C11	39498
4691	1/28/98	508	RTA00000187AF.i.14.2	M00001679B:H07	19406
4691	2/24/98	928	RTA00000340F.m.04.1	M00001679B:H07	19406
4692	2/24/98	10	RTA00000350R.c.12.1	M00001550D:A04	9728
4693	1/28/98	553	RTA00000201F.b.22.1	M00004344B:H04	35728
4694	1/28/98	459	RTA00000179AF.c.22.3	M00001393B:B09	22515
4694	1/28/98	351	RTA00000179AF.c.22.1	M00001393B:B09	22515
4695	1/28/98	351	RTA00000179AF.c.22.1	M00001393B:B09	22515
4695	1/28/98	459	RTA00000179AF.c.22.3	M00001393B:B09	22515
4696	2/24/98	235	RTA00000126A.o.23.1	M00001551A:B10	6268
4697	2/24/98	942	RTA00000126A.n.6.2	M00001551A:D04	79917
4698	2/24/98	228	RTA00000411F.k.19.1	M00003852D:E08	64200
4699	1/28/98	638	RTA00000193AF.1.05.2	M00004348A:A02	2815
4700	3/24/98	431	RTA00000425F.1.09.1	M00001638A:B04	75251
4701	1/28/98	540	RTA00000179AF.b.10.3	M00001391D:D10	0
4702	2/24/98	390	RTA00000355R.a.14.1	M00004187D:G09	10207
4703	1/28/98	429	RTA00000196F.i.19.1	M00001390C:C11	39498
4703	2/24/98	925	RTA00000353R.h.10.1	M00001390C:C11	39498
4704	2/24/98	930	RTA00000127A.h.22.2	M00001554A:E04	13155
4705	2/24/98	1193	RTA00000411F.k.14.1	M00003851A:C10	63987
4706	1/28/98	694	RTA00000201R.c.19.1	M00004370A:G05	22357
4707	3/24/98	80	RTA00000425F.p.12.1	M00001638C:G01	73219
4708	3/24/98	344	RTA00000425F.p.15.1	M00001638C:H07	31680
4709	1/28/98	743	RTA00000178AF.k.9.1	M00001381B:F06	16342
4710	2/24/98	202	RTA00000419F.g.22.1	M00003845D:A09	64515
4711	1/28/98	749	RTA00000178AR.i.13.4	M00001377B:H01	0
4712	3/24/98	217	RTA00000425F.j.16.1	M00001639D:F02	75631
4713	3/24/98	448	RTA00000425F.j.18.1	M00001639D:G12	75561
4714	1/28/98	385	RTA00000201F.c.24.1	M00004374D:E10	35731
4715	2/24/98	904	RTA00000127A.e.6.1	M00001553A:E07	5885
4716	2/24/98	620	RTA00000420F.a.07.1	M00004072C:F08	63405
4717	1/28/98	396	RTA00000179AR.b.02.3	M00001391B:G12	0
4718	2/24/98	463	RTA00000403F.o.22.1	M00001583A:D01	25076
4718	2/24/98	1084	RTA00000403F.o.22.2	M00001583A:D01	25076
4719	2/24/98	1225	RTA00000346F.j.13.1	M00003841C:E04	5337
4720	2/24/98	1221	RTA00000403F.o.17.1	M00001582D:A02	23085
4721	2/24/98	878	RTA00000413F.c.12.1	M00004083B:G03	65334
4722	2/24/98	764	RTA00000413F.c.17.1	M00004085B:B05	36831

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
4723	2/24/98	398	RTA00000407F.b.04.1	M00004086D:B09	63221
4724	2/24/98	1179	RTA00000341F.o.12.1	M00004144A:F04	2883
4725	2/24/98	7	RTA00000413F.d.12.1	M00004088C:A12	66467
4726	2/24/98	881	RTA00000413F.d.15.1	M00004088C:E04	64943
4727	2/24/98	463	RTA00000403F.o.22.1	M00001583A:D01	25076
4727	2/24/98	1084	RTA00000403F.o.22.2	M00001583A:D01	25076
4728	2/24/98	1084	RTA00000403F.o.22.2	M00001583A:D01	25076
4728	2/24/98	463	RTA00000403F.o.22.1	M00001583A:D01	25076
4729	1/28/98	506	RTA00000198R.o.09.1	M00003751B:A05	4310
4729	1/28/98	497	RTA00000198AF.o.09.1	M00003751B:A05	4310
4730	2/24/98	1015	RTA00000129A.b.6.2	M00001582A:H01	39111
4731	2/24/98	866	RTA00000407F.b.08.1	M00004088D:B03	37513
4732	2/24/98	943	RTA00000413F.c.03.1	M00004081D:H09	64527
4733	2/24/98	463	RTA00000403F.o.22.1	M00001583A:D01	25076
4733	2/24/98	1084	RTA00000403F.o.22.2	M00001583A:D01	25076
4734	1/28/98	490	RTA00000198AF.n.05.1	M00001687A:G01	24157
4735	2/24/98	798	RTA00000420F.c.04.1	M00004089A:B08	65007
4736	2/24/98	850	RTA00000420F.c.07.1	M00004089A:E02	65555
4737	2/24/98	199	RTA00000403F.m.13.2	M00001575D:A10	39077
4738	1/28/98	424	RTA00000187AR.j.24.1	M00001679D:B05	78356
4738	1/28/98	418	RTA00000187AR.k.01.1	M00001679D:B05	78356
4739	1/28/98	424	RTA00000187AR.j.24.1	M00001679D:B05	78356
4739	1/28/98	418	RTA00000187AR.k.01.1	M00001679D:B05	78356
4740	1/28/98	418	RTA00000187AR.k.01.1	M00001679D:B05	78356
4740	1/28/98	424	RTA00000187AR.j.24.1	M00001679D:B05	78356
4741	1/28/98	424	RTA00000187AR.j.24.1	M00001679D:B05	78356
4741	1/28/98	418	RTA00000187AR.k.01.1	M00001679D:B05	78356
4742	1/28/98	482	RTA00000187AF.j.7.1	M00001679C:F01	78091
4743	1/28/98	693	RTA00000198F.m.12.1	M00001679C:D05	4
4744	3/24/98	23	RTA00000522F.p.07.1	M00001670A:C11	76888
4745	1/28/98	497	RTA00000198AF.o.09.1	M00003751B:A05	4310
4745	1/28/98	506	RTA00000198R.o.09.1	M00003751B:A05	4310
4746	1/28/98	642	RTA00000189AF.i.14.1	M00003868B:G11	0
4747	2/24/98	1119	RTA00000126A.k.24.1	M00001550A:F07	39428
4748	2/24/98	654	RTA00000421F.a.05.1	M00001570C:G06	5278
4749	2/24/98	1146	RTA00000347F.h.02.1	M00004072D:H12	562
4750	2/24/98	137	RTA00000339R.a.06.1	M00001346A:E04	58694
4751	2/24/98	729	RTA00000403F.m.20.1	M00001576A:F11	707
4751	2/24/98	437	RTA00000403F.m.20.2	M00001576A:F11	707
4752	2/24/98	627	RTA00000408F.p.21.1	M00001579A:C03	77930
4753	2/24/98	735 535	RTA00000420F.a.11.1	M00004073C:D04	66460
4754	2/24/98	525	RTA00000348R.d.24.1	M00001349B:G05	5774
4755	2/24/98	624	RTA00000420F.a.16.1	M00004075D:C10	63345
4756	2/24/98	437	RTA00000403F.m.20.2	M00001576A:F11	707
4756	2/24/98	729	RTA00000403F.m.20.1	M00001576A:F11	707
4757	2/24/98	437	RTA00000403F.m.20.2	M00001576A:F11	707
4757	2/24/98	729	RTA00000403F.m.20.1	M00001576A:F11	707
4758	1/28/98	499	RTA00000199F.b.22.2	M00003791C:E09	17018
4759	2/24/98	843	RTA00000418F.g.03.1	M00001579C:E06	78737
4760	2/24/98	956	RTA00000423F.I.06.1	M00004062A:H06	38136

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority Appln			
4761	Appln 2/24/98	826	RTA00000422F.e.07.1	M00001579C:G05	38964
4761	2/24/98	832	RTA00000422F.c.07.1 RTA00000403F.o.10.2	M00001579C:G05	38964
	2/24/98	826	RTA00000403F.6.10.2	M00001579C:G05	38964
4762	2/24/98	832	RTA00000422F.c.07.1 RTA00000403F.o.10.2	M00001579C:G05	38964
4762		826	RTA00000403F.6.10.2 RTA00000422F.e.07.1	M00001579C:G05	38964
4763	2/24/98	832	RTA00000422F.c.07.1 RTA00000403F.o.10.2	M00001579C:G05	38964
4763	2/24/98		RTA00000403F.0.10.2 RTA00000422F.e.07.1	M00001579C:G05	38964
4764	2/24/98	826 832	RTA00000422F.c.07.1 RTA00000403F.o.10.2	M00001579C:G05	38964
4764	2/24/98		RTA00000403F.b.18.1	M00001379C:G03	39873
4765	2/24/98	1159		M00004078C:104 M00003839D:E02	64679
4766	2/24/98	1122	RTA00000419F.f.16.1	M00003839D.E02 M00004080A:F01	65117
4767	2/24/98	1053	RTA00000413F.b.24.1	M00004080A:F01	64013
4768	2/24/98	1052	RTA00000420F.b.02.1		4022
4769	2/24/98	157	RTA00000339F.a.23.1	M00001361B:C07	6707
4770	1/28/98	452	RTA00000199F.d.19.2	M00003813D:H12	
4771	2/24/98	480	RTA00000411F.i.15.1	M00003837C:G08	31612
4772	2/24/98	125	RTA00000403F.m.18.1	M00001576A:B09	39185
4773	2/24/98	548	RTA00000413F.b.12.1	M00004077B:H11	64932
4774	2/24/98	814	RTA00000408F.1.24.1	M00001530B:G09	34263
4775	1/28/98	688	RTA00000193AF.g.3.1	M00004050D:A06	5567
4776	1/28/98	451	RTA00000200AF.b.20.1	M00004043A:D02	40403
4777	1/28/98	456	RTA00000200AF.b.12.1	M00004040B:F10	22053
4778	2/24/98	849	RTA00000122A.n.16.1	M00001517A:G08	80553
4779	1/28/98	12	RTA00000183AF.i.15.2	M00001529B:C04	2642
4779	2/24/98	379	RTA00000349R.j.07.1	M00001529B:C04	2642
4780	1/28/98	12	RTA00000183AF.i.15.2	M00001529B:C04	2642
4780	2/24/98	379	RTA00000349R.j.07.1	M00001529B:C04	2642
4781	1/28/98	512	RTA00000191AF.c.3.1	M00003987D:D06	3549
4782	2/24/98	431	RTA00000399F.j.15.1	M00001578C:G06	1261
4783	1/28/98	586	RTA00000199R.o.11.1	M00003976C:A-10	23172
4784	1/28/98	496	RTA00000190AF.p.3.1	M00003975B:F03	2378
4785	2/24/98	340	RTA00000408F.l.13.1	M00001530A:B12	4423
4786	1/28/98	617	RTA00000179AF.d.13.3	M00001394A:F01	6583
4787	2/24/98	779	RTA00000408F.1.16.1	M00001530A:F12	73468
4788	1/28/98	387	RTA00000191AF.j.14.1	M00004073A:H12	1002
4788	2/24/98	632	RTA00000191AF.j.14.1	M00004073A:H12	1002
4789	1/28/98	464	RTA00000199AF.I.14.1	M00003917A:D02	22865
4790	2/24/98	668	RTA00000408F.m.05.2	M00001530C:G10	23384
4791	2/24/98	1066	RTA00000123A.f.2.1	M00001531A:H03	80379
4792	2/24/98	213	RTA00000123A.f.3.1	M00001531A:H07	44017
4793	2/24/98	43	RTA00000420F.g.04.1	M00004891B:B12	0
4794	2/24/98	302	RTA00000356R.f.18.1	M00004692A:H10	0
4795	2/24/98	308	RTA00000353R.d.11.1	M00004692A:H08	0
4796	2/24/98	975	RTA00000411F.n.02.1	M00003870B:F04	78049
4797	1/28/98	722	RTA00000199R.j.24.1	M00003895C:A10	0
4798	2/24/98	643	RTA00000420F.1.14.2	M00005230D:F06	0
4799	1/28/98	480	RTA00000181AF.o.08.2	M00001457C:H12	849
4800	1/28/98	518	RTA00000199AF.n.22.1	M00003971A:A06	23064
4801	1/28/98	618	RTA00000192AF.a.14.1	M00004111D:A08	6874
4802	2/24/98	937	RTA00000121A.n.15.1	M00001511A:G08	40849
4803	2/24/98	821	RTA00000420F.h.16.1	M00004927A:E06	0

Applin		SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
A805 2/24/98 172			•	•			
4806 2/24/98 172	4	1804			RTA00000121A.n.2.1	M00001511A:A05	33585
4807 2/24/98 1224	4	1805	2/24/98	551	RTA00000340F.b.05.1	M00001513A:G07	0
4808 1/28/98 631 RTA00000200AF.h.19.2 M00001514A:B04 81366 4809 2/24/98 809 RTA00000426P.i.23.1 M00001514A:B08 81366 4811 2/24/98 809 RTA00000426P.i.23.1 M00005134A:B01 0 4811 2/24/98 650 RTA00000122A.g.17.1 M00005134A:B08 32655 4812 2/24/98 650 RTA00000010AF.c.16.1 M000005136D:D06 0 4813 1/28/98 425 RTA00000191AF.j.15.2 M000005136D:G06 0 4815 1/28/98 387 RTA00000191AF.j.14.1 M00004073A:H12 1002 4816 2/24/98 632 RTA0000012A.j.i8.1 M00004573A:H12 1002 4817 2/24/98 837 RTA0000012A.j.i8.1 M0000473A:H12 1002 4817 2/24/98 837 RTA0000012A.j.i8.1 M0000473A:H12 1002 4818 2/24/98 73 RTA000002AP.d.20.1 M0000487C:D03 0 4819 1/28/98 734 RTA00000	4	1806	2/24/98	172	RTA00000420F.i.17.1	M00005101C:B09	0
4809 2/24/98 80 RTA00000122A.g.16.1 M00001514A:B04 81366 4810 2/24/98 809 RTA00000420F.i.23.1 M000005134A:D11 0 4811 2/24/98 650 RTA000000420F.i.23.1 M000005134A:D11 0 4812 2/24/98 73 RTA00000413F.p.15.2 M00005136D:D06 0 4813 1/28/98 425 RTA00000013F.p.15.2 M00005136D:D06 0 4814 2/24/98 60 RTA00000013F.p.15.2 M00005136D:D06 0 4815 1/28/98 387 RTA00000127A.p.14.1 M00004073A:H12 1002 4815 2/24/98 632 RTA00000012A.p.14.1.1 M00004870:D05 0 4817 2/24/98 837 RTA00000012A.p.1.8.1 M00004973A:H12 1002 4817 2/24/98 73 RTA00000012A.p.1.8.1 M00004870:D05 81317 4820 1/28/98 734 RTA0000001A.p.1.2.1 M00004870:D05 81317 4821 1/28/98 432 RTA00000A.p.	4	1807	2/24/98	1224	RTA00000122A.h.24.1	M00001514A:A12	48
4810 2/24/98 809 RTA00000420F.i.23.1 M00005134A:D11 0 4811 2/24/98 650 RTA00000122A:g.17.1 M00005134A:D11 0 4813 1/28/98 425 RTA00000413F.p.15.2 M00005136D:D60 0 4813 1/28/98 425 RTA00000413F.p.17.2 M00005136D:D60 0 4814 2/24/98 60 RTA00000191AF.j.1.2 M00004073A:H12 1002 4815 1/28/98 387 RTA00000191AF.j.1.1 M00004073A:H12 1002 4816 2/24/98 837 RTA0000012A.j.18.1 M00004073A:H12 1002 4817 2/24/98 173 RTA0000012A.j.18.1 M00004087C:D60 0 4818 2/24/98 173 RTA0000012A.j.1.1 M0000487C:D60 0 4820 1/28/98 73 RTA0000020AF.d.20.1 M0000487A:G08 26600 4821 1/28/98 73 RTA0000020AF.d.20.1 M0000487A:G08 26600 4821 1/28/98 42 RTA00000141F.p.01	4	1808	1/28/98	631	RTA00000200AF.h.19.2	M00004151D:E03	0
4811 2/24/98 650 RTA00000122A.g.17.1 M00001514A:B08 32655 4812 2/24/98 73 RTA00000200AF.c.16.1 M00001516D:D06 0 4813 1/28/98 425 RTA00000200AF.c.16.1 M00004064D:A11 23433 4814 2/24/98 60 RTA00000191AF.j.14.1 M00004073A:H12 1002 4815 1/28/98 387 RTA00000191AF.j.14.1 M00004073A:H12 1002 4816 2/24/98 837 RTA0000012AF.j.14.1 M00004073A:H12 1002 4816 2/24/98 123 RTA0000012AF.j.14.1 M0000457C:D06 0 4817 2/24/98 123 RTA000002C0AF.d.21.1 M00005173B:F01 0 4819 1/28/98 73 RTA00000220AF.d.20.1 M00004087A:G08 26600 4820 1/28/98 733 RTA00000220AF.d.20.1 M0000408TA:G08 26600 4821 1/28/98 442 RTA0000019TAF.d.20.1 M0000408TA:G08 26600 4822 1/28/98 457 <	2	1809	2/24/98	80	RTA00000122A.g.16.1	M00001514A:B04	81366
4812 2/24/98 73 RTA00000413F.p.I5.2 M00005136D:D06 0 4813 1/28/98 425 RTA00000200AF.c.16.1 M00004064D:A11 23433 4814 2/24/98 60 RTA00000191AF.j.I4.1 M00004073A:H12 1002 4815 1/28/98 387 RTA00000191AF.j.I4.1 M00004073A:H12 1002 4816 2/24/98 632 RTA00000122A.j.I8.1 M00004897C:D06 0 4817 2/24/98 123 RTA00000122A.j.I8.1 M00001516A:D05 81317 4818 2/24/98 734 RTA0000012AP.j.22.1 M0000487C:D03 0 4820 1/28/98 733 RTA0000020AF.d.20.1 M0000487A:G08 26600 4821 1/28/98 442 RTA00000191AF.I.9.1 M000048TA:G08 26600 4822 1/28/98 457 RTA00000197AF.e.02.1 M000048TC:D03 0 4823 1/28/98 457 RTA00000197AF.e.24.1 M00003875A:C04 73308 4825 2/24/98 839 RTA00	4	1810	2/24/98	809	RTA00000420F.i.23.1	M00005134A:D11	0
4813 1/28/98 425 RTA00000200AF.c.16.1 M00004064D:A11 23433 4814 2/24/98 60 RTA00000191AF.j.14.1 M00004073A:H12 1002 4815 1/28/98 387 RTA00000191AF.j.14.1 M00004073A:H12 1002 4816 2/24/98 632 RTA00000122A.j.18.1 M00001516A:D05 81317 4817 2/24/98 77 RTA00000122A.j.18.1 M00001516A:D05 81317 4818 2/24/98 77 RTA0000022A.j.18.1 M00004087C:D03 0 4819 1/28/98 734 RTA0000020AF.d.20.1 M00004087A:G08 26600 4820 1/28/98 733 RTA0000020AF.d.20.1 M00004087C:D03 0 4821 1/28/98 442 RTA00000191AF.19.1 M00004087C:D03 0 4822 1/28/98 457 RTA00000191AF.19.1 M00004081C:D12 14391 4824 1/28/98 457 RTA00000191AF.19.1 M00004081C:D12 14391 4825 2/24/98 852 RTA0	4	1811	2/24/98	650	RTA00000122A.g.17.1	M00001514A:B08	32655
4814 2/24/98 60 RTA00000413F.p.17.2 M00005136D:G06 0 4815 1/28/98 387 RTA00000191AF.j.14.1 M00004073A:H12 1002 4816 2/24/98 837 RTA00000191AF.j.14.1 M00004073A:H12 1002 4817 2/24/98 837 RTA00000120A.j.18.1 M00001516A:D05 81317 4818 2/24/98 77 RTA00000200AF.d.20.1 M00004087C:D03 0 4819 1/28/98 734 RTA00000200AF.d.20.1 M00004087A:G08 26600 4820 1/28/98 733 RTA00000200AF.d.20.1 M00004087A:G08 26600 4821 1/28/98 442 RTA00000191AF.l.9.1 M00004087A:G08 26600 4821 1/28/98 457 RTA00000191AF.l.9.1 M00004081C:D12 14391 4824 1/28/98 457 RTA00000191AF.l.9.1 M00004081C:D12 14391 4824 1/28/98 457 RTA00000141F.b.01.1 M000003875A:C04 73308 4825 1/28/98 674	4	1812	2/24/98	73	RTA00000413F.p.15.2	M00005136D:D06	0
4815 1/28/98 387 RTA00000191AF.j.14.1 M00004073A:H12 1002 4816 2/24/98 632 RTA000004074D.h.01.1 M000004073A:H12 1002 4816 2/24/98 123 RTA00000420F.h.01.1 M00004073A:H12 1002 4817 2/24/98 77 RTA00000420F.h.01.1 M00001516A:D05 81317 4818 2/24/98 734 RTA00000200AF.d.21.1 M00004087C:D03 0 4820 1/28/98 733 RTA00000200AF.d.21.1 M00004087C:D03 0 4821 2/24/98 1003 RTA00000420F.k.08.2 M00005176C:C09 0 4822 1/28/98 442 RTA00000191AF.1.9.1 M00004081C:D12 14391 4824 2/24/98 552 RTA00000411F.n.12.1 M00003875A:C04 73308 4825 2/24/98 782 RTA00000414F.b.03.1 M00003871C:B05 40822 4827 1/28/98 674 RTA00000197AF.e.24.1 M00001456B:F10 39250 4828 1/28/98 718	4	1813	1/28/98	425	RTA00000200AF.c.16.1	M00004064D:A11	23433
## ## ## ## ## ## ## ## ## ## ## ## ##	4	1814	2/24/98	60	RTA00000413F.p.17.2	M00005136D:G06	0
4816 2/24/98 837 RTA00000420F.h.01.1 M00004897C:D06 0 4817 2/24/98 123 RTA00000122A.j.18.1 M00001516A:D05 81317 4818 2/24/98 77 RTA00000420F.j.22.1 M00005173B:F01 0 4819 1/28/98 733 RTA00000200AF.d.20.1 M00004087A:G08 26600 4820 1/28/98 733 RTA00000200AF.d.20.1 M0000457C:C09 0 4821 1/28/98 442 RTA00000191AF.19.1 M0000481C:H06 0 4823 1/28/98 457 RTA00000191AF.19.1 M00003875A:C04 73308 4824 2/24/98 552 RTA00000411F.n.12.1 M00003871C:B05 40822 4824 2/24/98 552 RTA00000419F.k.03.1 M00003871C:B05 40822 4827 1/28/98 674 RTA00000197AF.e.24.1 M00001456B:F10 39250 4827 1/28/98 3 RTA00000197AF.e.24.1 M000014156B:F10 39250 4829 1/28/98 718 RTA0	4	18 15	1/28/98	387	RTA00000191AF.j.14.1	M00004073A:H12	1002
4817 2/24/98 123 RTA00000122A.j.18.1 M00001516A:D05 81317 4818 2/24/98 77 RTA00000200AF.d.2.1 M00005173B:F01 0 4819 1/28/98 734 RTA00000200AF.d.2.1 M00004087C:D03 0 4820 1/28/98 733 RTA00000200AF.d.2.1 M00004087A:G08 26600 4821 2/24/98 1003 RTA00000191AF.I.9.1 M00004081C:H06 0 4822 1/28/98 442 RTA00000191AR.I.7.2 M00004081C:H06 0 4823 1/28/98 457 RTA00000191AR.I.7.2 M00004081C:D12 14391 4824 2/24/98 552 RTA00000419F.k.03.1 M00003875A:C04 73308 4825 2/24/98 839 RTA00000197AR.e.24.1 M00001456B:F10 39250 4827 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 4828 1/28/98 689 RTA00000197AF.e.24.1 M000041456B:F10 39250 4829 1/28/98 718 <t< td=""><td>4</td><td>4815</td><td>2/24/98</td><td>632</td><td>RTA00000191AF.j.14.1</td><td>M00004073A:H12</td><td>1002</td></t<>	4	4815	2/24/98	632	RTA00000191AF.j.14.1	M00004073A:H12	1002
4818 2/24/98 77 RTA00000420F.j.22.1 M00005173B:F01 0 4819 1/28/98 734 RTA00000200AF.d.20.1 M00004087C:D03 0 4820 1/28/98 733 RTA00000200AF.d.20.1 M00004087A:G08 26600 4821 2/24/98 1003 RTA00000191AF.I.9.1 M00004081C:H06 0 4822 1/28/98 442 RTA00000191AR.I.7.2 M00004081C:H06 0 4823 1/28/98 457 RTA00000191AR.I.7.2 M00004081C:H06 0 4824 2/24/98 552 RTA0000041Fr.0.12.1 M00003875A:C04 73308 4825 2/24/98 782 RTA00000419Fk.0.3.1 M00003875A:C04 73308 4826 2/24/98 839 RTA00000197AR.e.24.1 M00001456B:F10 39250 4827 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 4828 1/28/98 69 RTA00000197AF.e.22.1 M00001456B:F10 39250 4828 1/28/98 718 RT	4	1816	2/24/98	837	RTA00000420F.h.01.1	M00004897C:D06	0
A819	4	1817	2/24/98	123	RTA00000122A.j.18.1	M00001516A:D05	81317
4820 1/28/98 733 RTA00000200AF.d.20.1 M00004087A:G08 26600 4821 2/24/98 1003 RTA00000420F.k.08.2 M00005176C:C09 0 4822 1/28/98 442 RTA00000191AF.1.9.1 M00004081C:D12 14391 4823 1/28/98 457 RTA00000191AR.1.7.2 M00004081C:D12 14391 4824 2/24/98 552 RTA0000041Fn.12.1 M00003871C:B05 40822 4825 2/24/98 782 RTA00000419F.k.03.1 M00003871C:B05 40822 4826 2/24/98 839 RTA00000197AF.e.24.1 M00001456B:F10 39250 4827 1/28/98 674 RTA00000197AF.e.24.1 M00001456B:F10 39250 4828 1/28/98 669 RTA00000197AF.e.24.1 M00001408B:G06 23144 4830 2/24/98 1259 RTA00000197AF.e.2.1 M0000420B:G06 23144 4830 2/24/98 17 RTA00000197AF.e.2.1 M00004249B:C08 36432 4831 1/28/98 717	4	1818	2/24/98	77	RTA00000420F.j.22.1	M00005173B:F01	0
4821 2/24/98 1003 RTA00000420F.k.08.2 M00005176C:C09 0 4822 1/28/98 442 RTA00000191AF.I.9.1 M00004081C:H06 0 4823 1/28/98 457 RTA00000191AF.I.9.1 M00004081C:D12 14391 4824 2/24/98 552 RTA00000419F.k.03.1 M00003875A:C04 73308 4825 2/24/98 839 RTA00000419F.k.03.1 M00003875A:C04 73308 4826 2/24/98 839 RTA00000197AF.e.24.1 M00001456B:F10 39250 4827 1/28/98 674 RTA00000192AF.e.24.1 M00001456B:F10 39250 4828 1/28/98 689 RTA00000192AF.e.24.1 M00001456B:F10 39250 4829 1/28/98 718 RTA00000192AF.e.21.1 M00001408B:G06 23144 4830 2/24/98 1259 RTA00000020F.o.10.2 M00004269B:C08 36432 4831 1/28/98 697 RTA000000125A.g.16.1 M00001454A:C09 21497 4833 1/28/98 409		4819	1/28/98	734	RTA00000200AF.d.21.1	M00004087C:D03	0
4822 1/28/98 442 RTA00000191AF.I.9.1 M00004081C:H06 0 4823 1/28/98 457 RTA00000191AR.I.7.2 M00004081C:D12 14391 4824 2/24/98 552 RTA00000411F.n.12.1 M00003875c:C04 73308 4825 2/24/98 782 RTA00000414F.b.01.1 M00003871C:B05 40822 4826 2/24/98 839 RTA00000197AR.e.24.1 M00001456B:F10 39250 4827 1/28/98 674 RTA00000197AF.e.24.1 M00001456B:F10 39250 4828 1/28/98 689 RTA00000197AF.e.24.1 M00001426B:F10 39250 4828 1/28/98 69 RTA00000192AF.e.21.1 M00001418B:G01 0 4830 2/24/98 1259 RTA00000420F.I.9.2 M00004269B:C08 36432 4831 1/28/98 717 RTA00000125A.g.16.1 M0000144A:C09 21497 4833 1/28/98 717 RTA00000180AF.d.21.1 M0000420F.B.06 0 4834 2/24/98 829	4	4820	1/28/98	733	RTA00000200AF.d.20.1	M00004087A:G08	26600
4822 1/28/98 442 RTA00000191AF.I.9.1 M00004081C:H06 0 4823 1/28/98 457 RTA00000191AR.I.7.2 M00004081C:D12 14391 4824 2/24/98 552 RTA00000411F.n.I.2.1 M00003873C:C04 73308 4825 2/24/98 839 RTA00000414F.b.03.1 M00003871C:B05 40822 4826 2/24/98 839 RTA00000197AR.e.24.1 M00001456B:F10 39250 4827 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 4828 1/28/98 669 RTA00000196F.I.14.2 M00001408B:G06 23144 4830 2/24/98 1259 RTA00000196F.I.19.2 M00005231A:H04 0 4831 1/28/98 717 RTA00000200F.o.10.2 M00004269B:C08 36432 4832 2/24/98 107 RTA00000125A.g.16.1 M00001544A:C09 21497 4833 1/28/98 697 RTA00000141F.m.11.1 M00004271B:B06 8526 4836 3/24/98 426	4	1821	2/24/98	1003	RTA00000420F.k.08.2	M00005176C:C09	0
4824 2/24/98 552 RTA00000411F.n.12.1 M00003875A:C04 73308 4825 2/24/98 782 RTA00000419F.k.03.1 M00003871C:B05 40822 4826 2/24/98 839 RTA000000197AF.e.24.1 M00001456B:F10 39250 4827 1/28/98 674 RTA00000197AF.e.24.1 M00001456B:F10 39250 4828 1/28/98 669 RTA00000197AF.e.24.1 M00001456B:F10 39250 4829 1/28/98 669 RTA00000192AF.c.2.1 M00004121B:G01 0 4830 2/24/98 1259 RTA00000196F.l.14.2 M00004269B:C08 36432 4831 1/28/98 717 RTA00000125A.g.16.1 M00001454B:C09 21497 4833 1/28/98 697 RTA00000125A.g.16.1 M0000424FB:D66 0 4834 2/24/98 829 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000044Fk.22.1 M00001614A:A04 73197 4837 2/24/98 874				442	RTA00000191AF.l.9.1	M00004081C:H06	0
4825 2/24/98 782 RTA00000419F.k.03.1 M00003871C:B05 40822 4826 2/24/98 839 RTA00000414F.b.01.1 M00005212B:A02 0 4827 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 4828 1/28/98 669 RTA00000192AF.c.2.1 M0000145B:G01 0 4829 1/28/98 718 RTA00000196F.i.14.2 M00001408B:G06 23144 4830 2/24/98 1259 RTA00000196F.i.19.2 M00005231A:H04 0 4831 1/28/98 717 RTA00000200F.o.10.2 M00004269B:C08 36432 4832 2/24/98 107 RTA00000125A.g.l6.1 M00001544A:C09 21497 4833 1/28/98 697 RTA0000013AF.e.21.1 M00004271B:B06 0 4834 2/24/98 829 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000424F.b.2.1 M00001614A:A04 73197 4837 2/24/98 874 RTA	4	4823	1/28/98	457	RTA00000191AR.I.7.2	M00004081C:D12	14391
4826 2/24/98 839 RTA000001414F.b.01.1 M00005212B:A02 0 4827 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 4827 1/28/98 3 RTA00000197AF.e.24.1 M00001412B:G01 0 4828 1/28/98 718 RTA00000196F.l.14.2 M00001412B:G01 0 4829 1/28/98 718 RTA00000196F.l.14.2 M0000143B:G06 23144 4830 2/24/98 1259 RTA00000420F.l.19.2 M00004269B:C08 36432 4831 1/28/98 717 RTA00000200F.o.10.2 M00004269B:C08 36432 4832 2/24/98 107 RTA00000125A.g.16.1 M00001544A:C09 21497 4833 1/28/98 697 RTA000000193AF.e.21.1 M0000427B:B06 0 4834 2/24/98 829 RTA00000041F.m.11.1 M00001614A:A04 73196 4835 1/28/98 409 RTA00000042F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874	4	1824	2/24/98	552	RTA00000411F.n.12.1	M00003875A:C04	73308
4827 1/28/98 674 RTA00000197AR.e.24.1 M00001456B:F10 39250 4827 1/28/98 3 RTA00000197AF.e.24.1 M00001456B:F10 39250 4828 1/28/98 669 RTA00000192AF.c.2.1 M00004121B:G01 0 4829 1/28/98 718 RTA00000196F.l.14.2 M00001408B:G06 23144 4830 2/24/98 1259 RTA00000420F.l.19.2 M00002331A:H04 0 4831 1/28/98 717 RTA00000200F.o.10.2 M00004269B:C08 36432 4832 2/24/98 107 RTA00000125A.g.16.1 M00001544A:C09 21497 4833 1/28/98 697 RTA00000193AF.e.21.1 M00004271B:B06 0 4834 2/24/98 829 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 409 RTA000000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000424F.k.21.1 M00001614A:A04 73197 4838 3/24/98 136	4	1825	2/24/98	782	RTA00000419F.k.03.1	M00003871C:B05	40822
4827 1/28/98 3 RTA00000197AF.e.24.1 M00001456B:F10 39250 4828 1/28/98 669 RTA00000192AF.c.2.1 M00004121B:G01 0 4829 1/28/98 718 RTA00000196F.i.14.2 M00005231A:H04 0 4830 2/24/98 1259 RTA00000200F.o.10.2 M00002531A:H04 0 4831 1/28/98 717 RTA00000200F.o.10.2 M00002469B:C08 36432 4832 2/24/98 107 RTA00000125A.g.16.1 M00001544A:C09 21497 4833 1/28/98 697 RTA0000013AF.e.21.1 M00004271B:B06 0 4834 2/24/98 829 RTA0000013AF.e.21.1 M00001418D:B06 8526 4835 1/28/98 409 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000424F.k.21.1 M00001614C:E11 72943 4839 2/24/98 636 RTA0	4	4826	2/24/98	839	RTA00000414F.b.01.1	M00005212B:A02	0
4828 1/28/98 669 RTA00000192AF.c.2.1 M00004121B:G01 0 4829 1/28/98 718 RTA00000196F.l.14.2 M00001408B:G06 23144 4830 2/24/98 1259 RTA00000420F.l.19.2 M00005231A:H04 0 4831 1/28/98 717 RTA00000200F.o.10.2 M00004269B:C08 36432 4832 2/24/98 107 RTA00000125A.g.16.1 M00001544A:C09 21497 4833 1/28/98 697 RTA0000013AF.e.21.1 M00004271B:B06 0 4834 2/24/98 829 RTA00000180AF.d.1.3 M00001418D:B06 8526 4835 1/28/98 409 RTA00000424F.k.21.1 M00001418D:B06 8526 4836 3/24/98 426 RTA00000346F.o.22.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000424F.m.22.1 M00001614C:E11 72943 4839 2/24/98 636 RTA00000418F.e.21.1 M0000157B:A03 74773 4840 2/24/98 120	4	4827	1/28/98	674	RTA00000197AR.e.24.1	M00001456B:F10	39250
4829 1/28/98 718 RTA00000196F.I.14.2 M00001408B:G06 23144 4830 2/24/98 1259 RTA00000420F.I.19.2 M00005231A:H04 0 4831 1/28/98 717 RTA00000200F.o.10.2 M00004269B:C08 36432 4832 2/24/98 107 RTA00000125A.g.16.1 M00001544A:C09 21497 4833 1/28/98 697 RTA0000013AF.e.21.1 M00004271B:B06 0 4834 2/24/98 829 RTA0000011F.m.11.1 M00003867A:D12 73196 4835 1/28/98 409 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000346F.o.22.1 M00001614C:E11 72943 4839 2/24/98 636 RTA00000424F.m.22.1 M00001577B:A03 74773 4840 2/24/98 1202 RTA00000347F.h.10.1 M00001542A:E04 80619 4841 2/24/98 221	4	482 7	1/28/98	3	RTA00000197AF.e.24.1	M00001456B:F10	39250
4830 2/24/98 1259 RTA00000420F.I.19.2 M00005231A:H04 0 4831 1/28/98 717 RTA00000200F.o.10.2 M00004269B:C08 36432 4832 2/24/98 107 RTA00000125A.g.16.1 M00001544A:C09 21497 4833 1/28/98 697 RTA00000193AF.e.21.1 M00004271B:B06 0 4834 2/24/98 829 RTA00000411F.m.11.1 M00003867A:D12 73196 4835 1/28/98 409 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000346F.o.22.1 M00001614C:E11 72943 4838 3/24/98 136 RTA000000424F.m.22.1 M0000157B:A03 74773 4840 2/24/98 636 RTA00000347F.h.10.1 M0000157B:A03 74773 4841 2/24/98 1030 RTA00000347F.h.10.1 M00001546B:F12 11500 4843 2/24/98 221	4	4828	1/28/98	669	RTA00000192AF.c.2.1	M00004121B:G01	0
4831 1/28/98 717 RTA00000200F.o.10.2 M00004269B:C08 36432 4832 2/24/98 107 RTA00000125A.g.16.1 M00001544A:C09 21497 4833 1/28/98 697 RTA00000193AF.e.21.1 M00004271B:B06 0 4834 2/24/98 829 RTA00000411F.m.11.1 M00003867A:D12 73196 4835 1/28/98 409 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000346F.o.22.1 M00004300C:H09 7381 4838 3/24/98 136 RTA00000418F.e.21.1 M00001577B:A03 74773 4840 2/24/98 636 RTA00000347F.h.10.1 M00001577B:A03 74773 4841 2/24/98 1030 RTA00000347F.h.10.1 M00001542A:E04 80619 4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 193		4829	1/28/98	718	RTA00000196F.l.14.2	M00001408B:G06	23144
4832 2/24/98 107 RTA00000125A.g.16.1 M00001544A:C09 21497 4833 1/28/98 697 RTA00000193AF.e.21.1 M00004271B:B06 0 4834 2/24/98 829 RTA00000411F.m.11.1 M00003867A:D12 73196 4835 1/28/98 409 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000346F.o.22.1 M00004300C:H09 7381 4838 3/24/98 136 RTA00000424F.m.22.1 M00001614C:E11 72943 4839 2/24/98 636 RTA00000418F.e.21.1 M00001577B:A03 74773 4840 2/24/98 1202 RTA00000347F.h.10.1 M00004206A:E02 22779 4841 2/24/98 1030 RTA00000125A.c.17.1 M00001546B:F12 11500 4842 2/24/98 753 RTA00000347F.b.10.1 M00001546C:C07 8044 4844 2/24/98 193		4830	2/24/98	1259	RTA00000420F.1.19.2	M00005231A:H04	0
4833 1/28/98 697 RTA00000193AF.e.21.1 M00004271B:B06 0 4834 2/24/98 829 RTA00000411F.m.11.1 M00003867A:D12 73196 4835 1/28/98 409 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000346F.o.22.1 M00001614C:E11 72943 4838 3/24/98 136 RTA00000424F.m.22.1 M00001577B:A03 74773 4840 2/24/98 636 RTA00000347F.h.10.1 M00001577B:A03 74773 4841 2/24/98 1202 RTA00000347F.h.10.1 M0000154CB:E02 22779 4841 2/24/98 1030 RTA00000125A.c.17.1 M0000154CB:F12 11500 4842 2/24/98 753 RTA00000345F.o.13.1 M0000154GB:F12 11500 4843 2/24/98 193 RTA00000347F.b.10.1 M00001547A:F06 0 4844 2/24/98 1177		4831	1/28/98	717		M00004269B:C08	36432
4834 2/24/98 829 RTA00000411F.m.11.1 M00003867A:D12 73196 4835 1/28/98 409 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000346F.o.22.1 M00004300C:H09 7381 4838 3/24/98 136 RTA00000424F.m.22.1 M00001614C:E11 72943 4839 2/24/98 636 RTA00000418F.e.21.1 M00001577B:A03 74773 4840 2/24/98 1202 RTA00000347F.h.10.1 M00001540A:E02 22779 4841 2/24/98 1030 RTA00000125A.c.17.1 M00001540A:E04 80619 4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 221 RTA00000347F.b.10.1 M00001546C:C07 8044 4844 2/24/98 193 RTA00000126A.b.10.1 M00001547A:F06 0 4846 2/24/98 1177		4832	2/24/98	107	RTA00000125A.g.16.1	M00001544A:C09	21497
4835 1/28/98 409 RTA00000180AF.d.1.3 M00001418D:B06 8526 4836 3/24/98 426 RTA00000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000346F.o.22.1 M00004300C:H09 7381 4838 3/24/98 136 RTA00000424F.m.22.1 M00001614C:E11 72943 4839 2/24/98 636 RTA00000418F.e.21.1 M00001577B:A03 74773 4840 2/24/98 1202 RTA00000347F.h.10.1 M0000154C:E02 22779 4841 2/24/98 1030 RTA00000125A.c.17.1 M0000154AB:F02 11500 4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 221 RTA00000347F.b.10.1 M00001546C:C07 8044 4845 2/24/98 193 RTA00000126A.b.10.1 M00001547A:F11 81279 4846 2/24/98 1177 RTA00000126A.b.9.1 M00001547A:F11 81279 4849 2/24/98 98	4	4833	1/28/98	697	RTA00000193AF.e.21.1	M00004271B:B06	
4836 3/24/98 426 RTA00000424F.k.21.1 M00001614A:A04 73197 4837 2/24/98 874 RTA00000346F.o.22.1 M00004300C:H09 7381 4838 3/24/98 136 RTA00000424F.m.22.1 M00001614C:E11 72943 4839 2/24/98 636 RTA00000418F.e.21.1 M00001577B:A03 74773 4840 2/24/98 1202 RTA00000347F.h.10.1 M00004206A:E02 22779 4841 2/24/98 1030 RTA00000125A.c.17.1 M00001542A:E04 80619 4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 221 RTA00000414F.f.13.1 M00001546C:C07 8044 4844 2/24/98 193 RTA00000126A.b.10.1 M00001547A:F06 0 4846 2/24/98 1104 RTA00000126A.b.9.1 M00001547A:F01 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 625	4	4834	2/24/98	829	RTA00000411F.m.11.1	M00003867A:D12	73196
4837 2/24/98 874 RTA00000346F.o.22.1 M00004300C:H09 7381 4838 3/24/98 136 RTA00000424F.m.22.1 M00001614C:E11 72943 4839 2/24/98 636 RTA00000418F.e.21.1 M00001577B:A03 74773 4840 2/24/98 1202 RTA00000347F.h.10.1 M00004206A:E02 22779 4841 2/24/98 1030 RTA00000125A.c.17.1 M00001542A:E04 80619 4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 221 RTA00000414F.f.13.1 M00001546C:C07 8044 4844 2/24/98 193 RTA00000347F.b.10.1 M00001546C:C07 8044 4845 2/24/98 1104 RTA00000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000126A.d.19.1 M00001549A:F01 0 4850 1/28/98 710	4	4835	1/28/98				8526
4838 3/24/98 136 RTA00000424F.m.22.1 M00001614C:E11 72943 4839 2/24/98 636 RTA00000418F.e.21.1 M00001577B:A03 74773 4840 2/24/98 1202 RTA00000347F.h.10.1 M00004206A:E02 22779 4841 2/24/98 1030 RTA00000125A.c.17.1 M00001542A:E04 80619 4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 221 RTA00000414F.f.13.1 M00001546C:C07 8044 4844 2/24/98 193 RTA00000347F.b.10.1 M00001547A:F06 0 4845 2/24/98 1104 RTA00000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.b.9.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000126A.d.19.1 M00001549A:F01 0 4849 2/24/98 625 RTA00000126A.h.22.2 M00001549A:F01 0 4850 1/28/98 710 <td< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td></td<>							
4839 2/24/98 636 RTA00000418F.e.21.1 M00001577B:A03 74773 4840 2/24/98 1202 RTA00000347F.h.10.1 M00004206A:E02 22779 4841 2/24/98 1030 RTA00000125A.c.17.1 M00001542A:E04 80619 4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 221 RTA00000414F.f.13.1 M00005259D:H08 0 4844 2/24/98 193 RTA00000347F.b.10.1 M00001546C:C07 8044 4845 2/24/98 1104 RTA00000126A.b.10.1 M00001547A:F06 0 4846 2/24/98 1177 RTA00000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000126A.h.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.l.3.1 M00001549A:H11 40425							
4840 2/24/98 1202 RTA00000347F.h.10.1 M00004206A:E02 22779 4841 2/24/98 1030 RTA00000125A.c.17.1 M00001542A:E04 80619 4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 221 RTA00000414F.f.13.1 M00005259D:H08 0 4844 2/24/98 193 RTA00000347F.b.10.1 M00001546C:C07 8044 4845 2/24/98 1104 RTA00000126A.b.10.1 M00001547A:F06 0 4846 2/24/98 1177 RTA00000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA0000041F.l.03.1 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.l.3.1 M00001549A:H11 40425 4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							
4841 2/24/98 1030 RTA000000125A.c.17.1 M00001542A:E04 80619 4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 221 RTA00000414F.f.13.1 M00005259D:H08 0 4844 2/24/98 193 RTA00000347F.b.10.1 M00001546C:C07 8044 4845 2/24/98 1104 RTA00000126A.b.10.1 M00001547A:F06 0 4846 2/24/98 1177 RTA00000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000411F.l.03.1 M00003854D:A12 62702 4849 2/24/98 625 RTA00000126A.b.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.l.3.1 M00001549A:H11 40425							
4842 2/24/98 753 RTA00000345F.o.13.1 M00001546B:F12 11500 4843 2/24/98 221 RTA00000414F.f.13.1 M00005259D:H08 0 4844 2/24/98 193 RTA00000347F.b.10.1 M00001546C:C07 8044 4845 2/24/98 1104 RTA00000126A.b.10.1 M00001547A:F06 0 4846 2/24/98 1177 RTA00000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000411F.l.03.1 M00003854D:A12 62702 4849 2/24/98 625 RTA00000126A.b.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.l.3.1 M00001549A:H11 40425							
4843 2/24/98 221 RTA00000414F.f.13.1 M00005259D:H08 0 4844 2/24/98 193 RTA00000347F.b.10.1 M00001546C:C07 8044 4845 2/24/98 1104 RTA00000126A.b.10.1 M00001547A:F06 0 4846 2/24/98 1177 RTA00000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000411F.l.03.1 M00003854D:A12 62702 4849 2/24/98 625 RTA00000126A.b.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.l.3.1 M00001405B:D07 20864 4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							
4844 2/24/98 193 RTA00000347F.b.10.1 M00001546C:C07 8044 4845 2/24/98 1104 RTA00000126A.b.10.1 M00001547A:F06 0 4846 2/24/98 1177 RTA00000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000411F.l.03.1 M00003854D:A12 62702 4849 2/24/98 625 RTA00000126A.b.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.l.3.1 M00001405B:D07 20864 4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							
4845 2/24/98 1104 RTA000000126A.b.10.1 M00001547A:F06 0 4846 2/24/98 1177 RTA00000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000411F.I.03.1 M00003854D:A12 62702 4849 2/24/98 625 RTA00000126A.h.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.I.3.1 M00001405B:D07 20864 4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							
4846 2/24/98 1177 RTA000000126A.b.9.1 M00001547A:F11 81279 4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000411F.l.03.1 M000003854D:A12 62702 4849 2/24/98 625 RTA00000126A.h.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.l.3.1 M00001405B:D07 20864 4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							
4847 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 4848 2/24/98 98 RTA00000411F.l.03.1 M00003854D:A12 62702 4849 2/24/98 625 RTA00000126A.h.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.l.3.1 M00001405B:D07 20864 4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							
4848 2/24/98 98 RTA00000411F.I.03.1 M00003854D:A12 62702 4849 2/24/98 625 RTA00000126A.h.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.I.3.1 M00001405B:D07 20864 4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							
4849 2/24/98 625 RTA00000126A.h.22.2 M00001549A:F01 0 4850 1/28/98 710 RTA00000196AF.l.3.1 M00001405B:D07 20864 4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							
4850 1/28/98 710 RTA00000196AF.I.3.1 M00001405B:D07 20864 4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							
4851 2/24/98 1102 RTA00000126A.j.15.2 M00001549A:H11 40425							-
•							
	4	1681	2/24/98	1102	•	M00001549A:H11	40425

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NO.	Priority	Priority			
	Appln	Appln			
4852	3/24/98	467	RTA00000528F.h.02.2	M00001632C:D08	1701
4853	1/28/98	411	RTA00000179AF.j.13.3	M00001400B:H06	0
4854	2/24/98	1126	RTA00000136A.h.6.1	M00001550A:D09	81620
4855	1/28/98	712	RTA00000201F.b.21.1	M00004341B:G03	9071
4856	2/24/98	1257	RTA00000419F.h.21.1	M00003856C:B08	64828
4857	2/24/98	194	RTA00000124A.k.5.1	M00001538A:F12	80252
4858	1/28/98	508	RTA00000187AF.i.14.2	M00001679B:H07	19406
4858	2/24/98	928	RTA00000340F.m.04.1	M00001679B:H07	19406
4859	1/28/98	3	RTA00000197AF.e.24.1	M00001456B:F10	39250
4859	1/28/98	674	RTA00000197AR.e.24.1	M00001456B:F10	39250
4860	2/24/98	606	RTA00000420F.1.20.2	M00005232A:C10	0
4861	2/24/98	30	RTA00000411F.m.24.1	M00003870B:B08	77568
4862	2/24/98	31	RTA00000134A.j.10.1	M00001534A:G06	81383
4863	2/24/98	1136	RTA00000406F.c.05.1	M00003870A:H01	22077
4864	2/24/98	411	RTA00000420F.m.12.1	M00005234D:B04	0
4865	2/24/98	1086	RTA00000403F.n.22.2	M00001578B:B05	26775
4865	2/24/98	1085	RTA00000403F.n.22.1	M00001578B:B05	26775
4866	2/24/98	1018	RTA00000413F.j.21.1	M00004688A:A02	0
4867	2/24/98	657	RTA00000124A.k.20.1	M00001538A:C08	80913
4868	2/24/98	718	RTA00000124A.k.23.1	M00001538A:D03	81350
4869	2/24/98	1092	RTA00000125A.c.2.1	M00001542A:F06	40148
4870	2/24/98	615	RTA00000135A.b.23.1	M00001538A:D12	35241
4871	2/24/98	639	RTA00000414F.d.09.1	M00005231C:B01	0
4872	2/24/98	1086	RTA00000403F.n.22.2	M00001578B:B05	26775
4872	2/24/98	1085	RTA00000403F.n.22.1	M00001578B:B05	26775
4873	2/24/98	99	RTA00000420F.m.19.1	M00005254D:B08	0
4874	2/24/98	1085	RTA00000403F.n.22.1	M00001578B:B05	26775
4874	2/24/98	1086	RTA00000403F.n.22.2	M00001578B:B05	26775
4875	2/24/98	1086	RTA00000403F.n.22.2	M00001578B:B05	26775
4 8 75	2/24/98	1085	RTA00000403F.n.22.1	M00001578B:B05	26775
4876	1/28/98	725	RTA00000197AF.b.1.1	M00001441D:E04	12134
4877	2/24/98	215	RTA00000403F.j.18.1	M00001539D:E10	5790
4878	2/24/98	1010	RTA00000408F.n.16.2	M00001540C:B03	73720
4879	2/24/98	1074	RTA00000423F.h.11.1	M00003867C:E11	38977
4880	2/24/98	27	RTA00000420F.n.19.2	M00005259B:C01	0
4881	1/28/98	578	RTA00000180AF.g.17.1	M00001426A:A09	16653
4882	2/24/98	1172	RTA00000423F.h.03.1	M00003875D:D09	37903
4883	2/24/98	1008	RTA00000414F.f.07.1	M00005259C:B05	0
4884	2/24/98	582	RTA00000414F.e.14.1	M00005236B:F10	0
4885	1/28/98	178	RTA00000200F.o.03.1	M00004257C:H06	22807
4885	1/28/98	249	RTA00000200R.o.03.2	M00004257C:H06	22807
4885	1/28/98	85	RTA00000200R.o.03.1	M00004257C:H06	22807
4886	2/24/98	1223	RTA00000347F.a.14.1	M00001429D:F11	7421
4887	2/24/98	488	RTA00000339F.k.23.1	M00001429D:H12	0
4888	3/24/98	100	RTA00000424F.i.21.1	M00001596A:E07	73482
4889	3/24/98	64	RTA00000424F.i.24.1	M00001596A:G06	79101
4890	3/24/98	207	RTA00000424F.j.07.1	M00001596B:C11	79211
4891	3/24/98	327	RTA00000424F.j.08.1	M00001596B:D09	73972
4892	3/24/98	349	RTA00000424F.j.09.1	M00001596B:H05	74387
4893	3/24/98	154	RTA00000522F.h.13.1	M00001596C:F09	40823

SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
	Appln	Appin			
4894	2/24/98	1252	RTA00000400F.g.08.1	M00001639A:C11	1275
4895	2/24/98	261	RTA00000341F.b.06.1	M00003794A:E12	17008
4896	1/28/98	312	RTA00000193AF.h.2.1	M0000379111212	3273
4897	1/28/98	590	RTA00000190AF.d.2.1	M00003906B:F12	2444
4898	1/28/98	213	RTA0000019071.d.2.1	M00003700D:F12	12514
4899	2/24/98	333	RTA000002001.0.04.1 RTA00000399F.f.11.1	M00004280D:C12	40167
4900	1/28/98	249	RTA000003991.1.11.1 RTA00000200R.o.03.2	M00001467C:F01	22807
4900	1/28/98	178	RTA00000200R.0.03.1	M00004257C:H06	22807
4900	1/28/98	85	RTA000002001.0.03.1	M00004257C:H06	22807
4900	1/28/98	85 85	RTA00000200R.o.03.1	M00004257C:H06	22807
4901	1/28/98	178	RTA00000200R.o.03.1	M00004257C:H06	22807
4901	1/28/98	249	RTA000002001.0.03.1 RTA00000200R.o.03.2	M00004257C:H06	22807
4901	1/28/98	85	RTA00000200R.o.03.1	M00004257C:H06	22807
4902	1/28/98	249	RTA00000200R.o.03.1 RTA00000200R.o.03.2	M00004257C:H06	22807
	1/28/98	178	RTA00000200F.o.03.1	M00004257C:H06	22807
4902		249	RTA00000200F.0.03.1 RTA00000200R.o.03.2	M00004257C:H06	22807
4903	1/28/98	249 85	RTA00000200R.o.03.1	M00004257C:H06	22807
4903	1/28/98	85 178	RTA00000200F.o.03.1	M00004257C:H06	22807
4903	1/28/98	249	RTA00000200F.0.03.1 RTA00000200R.o.03.2	M00004257C:H06	22807
4904	1/28/98 1/28/98	249 85	RTA00000200R.o.03.1	M00004257C:H06 M00004257C:H06	22807
4904		83 178	RTA00000200R.o.03.1	M00004257C:H06	22807
4904	1/28/98	178	RTA00000200F.6.03.1 RTA00000425F.f.04.1	M00004237C.H00 M00001607A:B06	24633
4905	3/24/98	169	RTA00000425F.f.05.1	M00001607A:D00	24033
4906	3/24/98	109 44	RTA00000423F.I.03.1 RTA00000418F.k.14.1	M00001607A.D10 M00001639A:H06	76133
4907	2/24/98		RTA00000418F.k.14.1 RTA00000419F.l.02.1	M00001039A.H00 M00003879A:C01	75736
4908	2/24/98	1204 748	RTA00000419F.1.02.1 RTA00000346F.f.11.1	M00003879A:C01 M00003793C:D09	38528
4909	2/24/98	74 0 4	RTA00000346F.I.11.1 RTA00000339F.i.20.1	M00003793C.D09 M00001438D:C06	4356
4910	2/24/98	93	RTA00000339F.1.20.1 RTA00000200F.o.11.1	M00001438D.C00 M00004270A:F11	0
4911 4912	1/28/98 1/28/98	435	RTA00000200F.0.11.1 RTA00000182AR.c.22.1	M00004270A.[11 M00001467A:D08	16283
4912	1/28/98	683	RTA00000182AR.c.22.1 RTA00000187AR.j.01.1	M00001407A:D08	79028
4913	3/24/98	469	RTA00000187Ak.j.01.1 RTA00000522F.e.20.1	M000015790B:H10	26770
4915	1/28/98	172	RTA000003221.6.20.1 RTA00000186AF.p.09.2	M00001550D:1110	6879
4916	2/24/98	806	RTA00000186A1.p.03.2	M00001033C:E04 M00001413B:H09	0075
4917	1/28/98	677	RTA000003431.1.00.1	M00001419B:H11	39554
4918	1/28/98	443	RTA00000197AR.i.17.1	M00001490A:E11	3516
4919	2/24/98	863	RTA0000019771R.1.17.17 RTA00000406F.p.08.1	M00004032C:B02	37573
4920	3/24/98	55	RTA00000528F.e.23.1	M00001593B:D10	19242
4921	2/24/98	1211	RTA00000399F.f.14.1	M00001487D:C11	11483
4922	1/28/98	609	RTA00000196AF.n.05.1	M00001418B:F07	12531
4922	2/24/98	1120	RTA00000353R.I.23.1	M00001418B:F07	12531
4923	1/28/98	609	RTA00000196AF.n.05.1	M00001418B:F07	12531
4923	2/24/98	1120	RTA00000353R.1.23.1	M00001418B:F07	12531
4924	3/24/98	474	RTA00000522F.h.05.1	M00001595C:H11	73358
4925	1/28/98	284	RTA00000199F.d.10.2	M00003808C:B05	22049
4925	2/24/98	816	RTA00000354R.n.04.1	M00003808C:B05	22049
4926	2/24/98	1112	RTA00000418F.c.05.1	M00001487B:F02	76475
4927	1/28/98	687	RTA00000197AF.g.4.1	M00001464B:B03	8821
4928	2/24/98	990	RTA00000121A.h.19.1	M00001471A:D04	80334
4929	1/28/98	696	RTA00000180AR.d.16.3	M00001419D:C10	11393
4929	2/24/98	1184	RTA00000345F.h.08.1	M00001419D:C10	11393

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
NO.	Priority	Priority			
	Appln	Appln			
4930	1/28/98	751	RTA00000179AF.c.4.3	M00001392D:B11	0
4931	1/28/98	696	RTA00000180AR.d.16.3	M00001419D:C10	11393
4931	2/24/98	1184	RTA00000345F.h.08.1	M00001419D:C10	11393
4932	1/28/98	162	RTA00000201F.e.15.1	M00004444B:D11	9960
4933	1/28/98	126	RTA00000201F.d.16.1	M00004388B:A08	0
4934	1/28/98	332	RTA00000193AR.n.04.3	M00004375C:D01	9850
4935	2/24/98	838	RTA00000408F.k.19.1	M00001487C:G03	77593
4936	2/24/98	113	RTA00000401F.e.02.1	M00003805B:C04	0
4937	3/24/98	278	RTA00000522F.g.21.1	M00001595C:A09	77310
4938	3/24/98	54	RTA00000425F.f.19.1	M00001653D:G07	32635
4939	2/24/98	976	RTA00000419F.b.17.1	M00003808D:D04	63261
4940	2/24/98	59	RTA00000346F.j.08.1	M00003879B:A06	39951
4941	2/24/98	151	RTA00000341F.c.21.1	M00003789C:F06	7899
4942	1/28/98	67	RTA00000200AF.g.07.1	M00004128B:G01	0
4943	2/24/98	324	RTA00000340F.p.17.1	M00003750C:H05	0
4944	2/24/98	476	RTA00000345F.h.01.1	M00001441B:D11	10834
4945	2/24/98	245	RTA00000195AF.d.20.1	M00004117A:D11	37574
4945	1/28/98	87	RTA00000195AF.d.20.1	M00004117A:D11	37574
4946	2/24/98	245	RTA00000195AF.d.20.1	M00004117A:D11	37574
4946	1/28/98	87	RTA00000195AF.d.20.1	M00004117A:D11	37574
4947	2/24/98	991	RTA00000419F.b.10.1	M00001694C:G04	78566
4948	2/24/98	14	RTA00000419F.b.09.1	M00001694C:F12	78128
4949	1/28/98	261	RTA00000192AF.a.24.1	M00004114C:F11	13183
4950	1/28/98	24	RTA00000200AF.f.11.1	M00004111D:D11	0
4951	3/24/98	408	RTA00000522F.o.10.1	M00001660D:E05	78798
4952	1/28/98	328	RTA00000200AF.g.09.1	M00004131B:H09	22785
4952	1/28/98	26	RTA00000200R.g.09.1	M00004131B:H09	22785
4953	2/24/98	861	RTA00000419F.b.06.1	M00001694B:B08	76728
4954	3/24/98	24	RTA00000522F.n.08.1	M00001656A:D10	76343
4955	2/24/98	760	RTA00000423F.d.04.1	M00001694A:B12	11307
4956	3/24/98	220	RTA00000522F.o.18.1	M00001669B:H06	76366
4957	2/24/98	279	RTA00000418F.i.21.1	M00001596D:E10	78728
4958	1/28/98	84	RTA00000191AF.h.14.1	M00004056B:D09	13553
4959	1/28/98	284	RTA00000199F.d.10.2	M00003808C:B05	22049
4959	2/24/98	816	RTA00000354R.n.04.1	M00003808C:B05	22049
4960	2/24/98	217	RTA00000399F.o.17.1	M00001599D:A09	1106
4961	1/28/98	287	RTA00000200AF.b.07.1	M00004039C:C01	17125
4961	1/28/98	173	RTA00000200AR.b.07.1	M00004039C:C01	17125
4962	1/28/98	287	RTA00000200AF.b.07.1	M00004039C:C01	17125
4962	1/28/98	173	RTA00000200AR.b.07.1	M00004039C:C01	17125
4963	1/28/98	287	RTA00000200AF.b.07.1	M00004039C:C01	17125
4963	1/28/98	173	RTA00000200AR.b.07.1	M00004039C:C01	17125
4964	1/28/98	287	RTA00000200AF.b.07.1	M00004039C:C01	17125
4964	1/28/98	173	RTA00000200AR.b.07.1	M00004039C:C01	17125
4965	3/24/98	464	RTA00000522F.p.18.1	M00001671A:H06	76376 73322
4966	3/24/98	453	RTA00000522F.p.22.1	M00001671B:F02	3055
4967	2/24/98	54	RTA00000399F.o.01.1	M00001595C:E01 M00003771B:E05	3033 39911
4968	2/24/98	1219	RTA00000347F.e.20.1	M00003771B:E03	39084
4969	2/24/98	825 364	RTA00000404F.k.22.2 RTA00000404F.k.22.1	M00001635D:C12	39084
4969	2/24/98	364	K I AUUUUU4U4F.K.22. I	1VIOUUU1033D.C12	37007

SEQ ID NO:	Filing Date of	SEQ ID NO: in	Sequence Name	Clone Name	Cluster ID
	Priority	Priority			
	Appln	Appln			
4970	1/28/98	241	RTA00000200AF.I.17.1	M00004217C:D03	12771
4970	1/28/98	151	RTA00000200R.I.17.1	M00004217C:D03	12771
4970	1/28/98	202	RTA00000200R.1.17.2	M00004217C:D03	12771
4971	1/28/98	241	RTA00000200AF.I.17.1	M00004217C:D03	12771
4971	1/28/98	151	RTA00000200R.I.17.1	M00004217C:D03	12771
4971	1/28/98	202	RTA00000200R.I.17.2	M00004217C:D03	12771
4972	1/28/98	241	RTA00000200AF.l.17.1	M00004217C:D03	12771
4972	1/28/98	202	RTA00000200R.1.17.2	M00004217C:D03	12771
4972	1/28/98	151	RTA00000200R.l.17.1	M00004217C:D03	12771
4973	1/28/98	241	RTA00000200AF.I.17.1	M00004217C:D03	12771
4973	1/28/98	202	RTA00000200R.l.17.2	M00004217C:D03	12771
4973	1/28/98	151	RTA00000200R.I.17.1	M00004217C:D03	12771
4974	1/28/98	241	RTA00000200AF.l.17.1	M00004217C:D03	12771
4974	1/28/98	202	RTA00000200R.1.17.2	M00004217C:D03	12771
4974	1/28/98	151	RTA00000200R.1.17.1	M00004217C:D03	12771
4975	1/28/98	241	RTA00000200AF.I.17.1	M00004217C:D03	12771
4975	1/28/98	151	RTA00000200R.1.17.1	M00004217C:D03	12771
4975	1/28/98	202	RTA00000200R.l.17.2	M00004217C:D03	12771
4976	1/28/98	241	RTA00000200AF.1.17.1	M00004217C:D03	12771
4976	1/28/98	151	RTA00000200R.I.17.1	M00004217C:D03	12771
4976	1/28/98	202	RTA00000200R.1.17.2	M00004217C:D03	12771
4977	1/28/98	241	RTA00000200AF.1.17.1	M00004217C:D03	12771
4977	1/28/98	151	RTA00000200R.1.17.1	M00004217C:D03	12771
4977	1/28/98	202	RTA00000200R.1.17.2	M00004217C:D03	12771
4978	1/28/98	241	RTA00000200AF.I.17.1	M00004217C:D03	12771
4978	1/28/98	151	RTA00000200R.1.17.1	M00004217C:D03	12771
4978	1/28/98	202	RTA00000200R.1.17.2	M00004217C:D03	12771
4979	1/28/98	366	RTA00000192AF.o.19.1	M00004208D:H08	3549
4980	1/28/98	328	RTA00000200AF.g.09.1	M00004131B:H09	22785
4980	1/28/98	26	RTA00000200R.g.09.1	M00004131B:H09	22785
4981	1/28/98	245	RTA00000200AF.k.7.1	M00004193C:G11	0
4982	2/24/98	1036	RTA00000339F.k.08.1	M00001439B:A10	8133
4983	2/24/98	72	RTA00000347F.a.08.1	M00001592C:G04	3135
4984	2/24/98	1163	RTA00000341F.b.14.1	M00003763A:C01	5992
4985	2/24/98	278	RTA00000404F.c.10.1	M00001593B:E11	23534
4986	1/28/98	250	RTA00000192AF.j.21.1	M00004176D:B12	2289
4987	2/24/98	511	RTA00000341F.b.13.1	M00003762B:H09	0
4988	1/28/98	27	RTA00000192AF.i.12.1	M00004169C:C12	5319
4989	2/24/98	416	RTA00000404F.c.19.1	M00001594A:D06	39026
4990	2/24/98	351	RTA00000340F.p.20.1	M00003752B:C02	17008
4991	1/28/98	215	RTA00000192AR.e.14.3	M00004142A:D08	3300
4992	1/28/98	163	RTA00000192AR.e.13.3	M00004142A:B12	9457
4993	1/28/98	318	RTA00000200AF.g.17.1	M00004138A:H09	0
4994	2/24/98	1105	RTA00000340F.p.18.1	M00003751C:A04	287
4995	2/24/98	1080	RTA00000351R.g.06.1	M00003771D:G05	0
4996	2/24/98	478	RTA00000418F.h.08.1	M00001589B:E07	76401
4997	2/24/98	584	RTA00000418F.d.22.1	M00001573B:C06	75324
4998	2/24/98	493	RTA00000129A.d.1.2	M00001587A:F05	80058
4999	2/24/98	402	RTA00000420F.e.16.1	M00004110A:E04	63639
5000	2/24/98	1006	RTA00000129A.e.14.1	M00001587A:F08	80053

Appln	SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
1/28/98 659						
5003 2/24/98 122 RTA0000420F.f.06.1 M00004115D:D08 64812 5004 2/24/98 245 RTA00000195AF.d.20.1 M00004117A:D11 37574 5004 1/28/98 87 RTA00000195AF.d.20.1 M00004117A:D11 37574 5005 1/28/98 87 RTA00000195AF.d.20.1 M00004117A:D11 37574 5006 2/24/98 720 RTA00000195AF.d.20.1 M00004117A:D11 37574 5006 2/24/98 687 RTA00000195AF.d.20.1 M00001587A:G06 80119 5007 2/24/98 687 RTA00000350R.g.10.1 M00001587A:G06 80119 5008 3/24/98 18 RTA00000186AF.d.3.1 M00001587A:G08 75283 5010 1/28/98 447 RTA00000186AR.e.07.4 M00001623D:G03 4175 5011 1/28/98 526 RTA00000185AF.d.20.1 M00001587A:H03 0 5011 1/28/98 526 RTA00000185AF.d.20.1 M00001589B:E12 9738 5013 1/28/98 530	5001					
5004 2/24/98 245 RTA00000195AF.d.20.1 M00004117A:D11 37574 5004 1/28/98 87 RTA00000195AF.d.20.1 M00004117A:D11 37574 5005 1/28/98 87 RTA00000195AF.d.20.1 M00004117A:D11 37574 5005 2/24/98 245 RTA00000195AF.d.20.1 M00004117A:D11 37574 5006 2/24/98 720 RTA00000129A.d.2.4 M00001587A:G06 80119 5007 2/24/98 687 RTA00000522F.e.16.1 M00001587A:G06 80119 5008 3/24/98 18 RTA00000186AR.e.07.4 M00001587A:H03 0 5010 1/28/98 447 RTA00000186AR.e.07.4 M0000158D3:C03 4175 5010 1/28/98 526 RTA00000185AF.e.20.1 M0000158D3:C03 4175 5011 1/28/98 526 RTA00000185AF.d.24.2 M0000158D3:A11 2385 5012 2/24/98 1 RTA00000421F.a.06.1 M0000158D3:A11 2385 5015 1/28/98 530	5002	1/28/98				
5004 1/28/98 87 RTA00000195AF.d.20.1 M00004117A:D11 37574 5005 1/28/98 87 RTA00000195AF.d.20.1 M00004117A:D11 37574 5005 2/24/98 245 RTA00000195AF.d.20.1 M000004117A:D11 37574 5006 2/24/98 720 RTA00000195AF.d.2.4 M00001587A:G06 80119 5007 2/24/98 687 RTA0000035QF.e.16.1 M00001587C:C10 9026 5008 3/24/98 18 RTA0000018AF.d.8.1 M00001587A:H03 75283 5009 1/28/98 447 RTA00000186AR.e.07.4 M00001587A:H03 0 5010 1/28/98 526 RTA00000185AF.e.20.1 M000015890:G03 4175 5011 1/28/98 526 RTA00000185AF.e.20.1 M000015890:G03 4175 5012 2/24/98 1 RTA0000044F.a.02.1 M00001589E12 9738 5013 1/28/98 530 RTA00000415A.A.d.2 M00001589D:C03 6539 5014 2/24/98 131 <t< td=""><td>5003</td><td>2/24/98</td><td>122</td><td>RTA00000420F.f.06.1</td><td></td><td></td></t<>	5003	2/24/98	122	RTA00000420F.f.06.1		
5005 1/28/98 87 RTA00000195AF.d.20.1 M00004117A:D11 37574 5005 2/24/98 245 RTA00000195AF.d.20.1 M00004117A:D11 37574 5006 2/24/98 687 RTA00000129A.d.2.4 M00001587A:G06 80119 5007 2/24/98 687 RTA00000350R.g.10.1 M00001587A:G06 80119 5008 3/24/98 18 RTA00000198AF.d.8.1 M00001587A:H03 0 5009 1/28/98 447 RTA00000198AF.d.8.1 M00001587A:H03 0 5010 1/28/98 554 RTA00000186AR.e.07.4 M00001582D:G03 4175 5011 1/28/98 526 RTA00000185AF.e.20.1 M00001585A:D06 5865 5012 2/24/98 10 RTA00000494F.a.02.1 M00001589B:E12 9738 5013 1/28/98 530 RTA000004185AF.d.11.2 M00001589C:A11 2385 5015 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 5016 1/28/98 626	5004	2/24/98		RTA00000195AF.d.20.1	M00004117A:D11	
5005 2/24/98 245 RTA00000195AF.d.20.1 M00004117A:D11 37574 5006 2/24/98 720 RTA00000129A.d.2.4 M00001587A:G06 80119 5007 2/24/98 687 RTA00000350R.g.10.1 M00001587C:C10 9026 5008 3/24/98 18 RTA00000522F.e.16.1 M00001590A:C08 75283 5009 1/28/98 447 RTA00000186AR.e.07.4 M00001623D:G03 4175 5010 1/28/98 554 RTA00000186AR.e.07.3 M00001623D:G03 4175 5011 1/28/98 526 RTA00000185AF.e.20.1 M00001585A:D06 5865 5012 2/24/98 1 RTA00000185AF.d.24.2 M00001585A:D06 5865 5013 1/28/98 530 RTA00000494F.a.02.1 M00001585A:D06 5865 5014 2/24/98 1096 RTA00000185AF.d.24.2 M00001589C:A11 2385 5015 1/28/98 626 RTA00000185AF.d.11.2 M00001579D:C03 6539 5016 1/28/98 626	5004	1/28/98		RTA00000195AF.d.20.1	M00004117A:D11	
5006 2/24/98 720 RTA00000129A.d.2.4 M00001587A:G06 80119 5007 2/24/98 687 RTA00000350R.g.10.1 M00001587C:C10 9026 5008 3/24/98 18 RTA00000522F.e.16.1 M00001590A:C08 75283 5009 1/28/98 447 RTA00000198AF.d.8.1 M00001587A:H03 0 5010 1/28/98 400 RTA00000186AR.e.07.3 M00001623D:G03 4175 5011 1/28/98 526 RTA00000185AF.e.20.1 M00001585A:D06 865 5012 2/24/98 1 RTA00000185AF.e.20.1 M00001585A:D06 865 5013 1/28/98 530 RTA00000185AF.e.24.2 M00001589E:E12 9738 5014 2/24/98 1096 RTA00000185AF.d.24.2 M00001589C:A11 2385 5015 1/28/98 626 RTA00000185AF.d.11.2 M00001579D:C03 6539 5016 1/28/98 626 RTA00000185AF.d.11.1 M00001579D:C03 6539 5017 2/24/98 131	5005	1/28/98	87	RTA00000195AF.d.20.1	M00004117A:D11	
5007 2/24/98 687 RTA00000350R.g.10.1 M00001587C:C10 9026 5008 3/24/98 18 RTA00000522F.e.16.1 M00001590A:C08 75283 5009 1/28/98 447 RTA00000198AF.d.8.1 M00001587A:H03 0 5010 1/28/98 544 RTA00000186AR.e.07.4 M00001623D:G03 4175 5010 1/28/98 400 RTA00000185AF.e.20.1 M00001583A:D06 5865 5011 1/28/98 526 RTA0000044F.a.02.1 M00001589B:E12 9738 5012 2/24/98 1 RTA0000044F.a.02.1 M00001589B:E12 9738 5013 1/28/98 530 RTA0000042F.a.02.1 M00001589B:E12 9738 5014 2/24/98 1996 RTA0000043F.A.d.11.2 M00001589B:E12 9738 5015 1/28/98 131 RTA00000185AF.d.11.2 M00001579D:C03 6539 5015 1/28/98 626 RTA00000185AR.d.11.1 M00001579D:C03 6539 5016 1/28/98 626 <th< td=""><td>5005</td><td>2/24/98</td><td>245</td><td>RTA00000195AF.d.20.1</td><td>M00004117A:D11</td><td>37574</td></th<>	5005	2/24/98	245	RTA00000195AF.d.20.1	M00004117A:D11	37574
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5039 2/24/98 1130 RTA00000354R.p.01.1 M00004104C:H12 0						
5040 2/24/98 710 RTA00000413F.g.24.1 M00004104D:A04 65481				-		
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5043	1/28/98	558	RTA00000186AR.h.14.1	M00001632D:H07	0
5045	2/24/98	155	RTA00000413F.h.13.1	M00004107A:D01	65190
5046	2/24/98	926	RTA00000399F.k.20.1	M00001585C:D10	3003
5047	2/24/98	1194	RTA00000420F.e.05.1	M00004107D:E12	63908
5047	1/28/98	400	RTA00000186AR.e.07.3	M00001623D:G03	4175
5048	1/28/98	554	RTA00000186AR.e.07.4	M00001623D:G03	4175
5049	2/24/98	570	RTA00000405F.n.13.1	M00003824A:G10	23810
5050	2/24/98	334	RTA00000408F.p.05.1	M00001575B:B02	9649
5051	2/24/98	1029	RTA00000411F.f.04.1	M00003813A:G04	64526
5052	3/24/98	134	RTA00000424F.c.14.3	M00001476D:A09	76614
5053	2/24/98	396	RTA00000406F.e.21.1	M00003877D:G05	9090
5054	3/24/98	230	RTA00000424F.g.14.1	M00001572A:B06	74879
5055	2/24/98	617	RTA00000423F.f.23.1	M00003816C:E09	15390
5056	2/24/98	5	RTA00000408F.o.12.2	M00001572A:A10	78578
5057	2/24/98	689	RTA00000419F.p.03.1	M00004035A:G10	1937
5058	3/24/98	273	RTA00000424F.a.02.4	M00001575A:D06	78806
5059	2/24/98	241	RTA00000339F.d.13.1	M00001395C:F11	0
5060	3/24/98	237	RTA00000522F.c.01.1	M00001576A:C11	74938
5061	1/28/98	745	RTA00000183AF.m.11.1	M00001536D:G02	8927
5062	1/28/98	408	RTA00000183AR.I.15.1	M00001535C:E01	39383
5063	2/24/98	464	RTA00000195AF.c.12.1	M00003818B:G12	37582
5063	1/28/98	300	RTA00000195AF.c.12.1	M00003818B:G12	37582
5064	1/28/98	647	RTA00000197F.m.11.1	M00001530B:D10	16488
5065	2/24/98	464	RTA00000195AF.c.12.1	M00003818B:G12	37582
5065	1/28/98	300	RTA00000195AF.c.12.1	M00003818B:G12	37582
5066	3/24/98	395	RTA00000522F.d.06.1	M00001578B:A02	74809
5067	2/24/98	516	RTA00000339F.f.20.1	M00001399A:C03	6494
5068	2/24/98	890	RTA00000418F.j.19.1	M00001634D:D02	78399
5069	2/24/98	435	RTA00000340F.b.02.1	M00001503C:G05	10185
5070	3/24/98	175	RTA00000528F.d.18.1	M00001582C:E01	2684
5071	2/24/98	168	RTA00000411F.e.22.1	M00003812B:D07	63638
5072	2/24/98	1071	RTA00000404F.k.18.2	M00001635A:C06	5475
5073	2/24/98	189	RTA00000347F.a.13.1	M00001402D:F02	22446
5074	2/24/98	825	RTA00000404F.k.22.2	M00001635D:C12	39084
5074	2/24/98	364	RTA00000404F.k.22.1	M00001635D:C12	39084
5075	3/24/98	25	RTA00000425F.c.06.1	M00001585D:D11	78041
5076	3/24/98	186	RTA00000425F.c.07.1	M00001585D:F03	76042
5077	3/24/98	208	RTA00000424F.m.10.1	M00001586C:E06	34251
5078	2/24/98	420	RTA00000422F.b.16.1	M00003813B:A11	17045
5079	3/24/98	103	RTA00000424F.b.22.1	M00001530A:F11	72971
5079	3/24/98	88	RTA00000424F.b.22.4	M00001530A:F11	72971
5080	3/24/98	318	RTA00000523F.a.01.1	M00001671C:F11	74923
5081	2/24/98	676	RTA00000411F.g.21.1	M00003823D:G05	64500
5082	3/24/98	3	RTA00000528F.b.23.1	M00001479C:F10	1605
5083	2/24/98	1244	RTA00000418F.h.23.1	M00001591A:B08	75153
5084	2/24/98	321	RTA00000339F.c.21.1	M00001389C:A08	5325
5085	1/28/98	429	RTA00000196F.i.19.1	M00001390C:C11	39498
5085	2/24/98	925	RTA00000353R.h.10.1	M00001390C:C11	39498
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S087 3/24/98 471	5086			RTA00000196F.i.19.1		
S088 2/24/98 103	5086	2/24/98	925	RTA00000353R.h.10.1	M00001390C:C11	
S089 2/24/98 1148 RTA00000345F.d.23.1 M00001390D:E03 5862	5087	3/24/98	471	RTA00000528F.c.11.1		
18267 1870	5088	2/24/98	103	RTA00000418F.j.12.1		
Sop	5089	2/24/98	1148	RTA00000345F.d.23.1		
1/28/98 661 RTA00000198R.b.04.1 M00001565A:H09 0	5090	2/24/98	87	RTA00000403F.1.20.1		
1975 1976 1977 1978 1977 1978 1979	5091	3/24/98	427	RTA00000522F.b.08.1	M00001570D:E06	
S094 2/24/98 1243 RTA00000404F.j.19.1 M00001631D:H10 O	5092	1/28/98	661	RTA00000198R.b.04.1	M00001565A:H09	-
1/28/98 750 RTA00000198AF.a.19.1 M00001561D:C05 0	5093	2/24/98	200	RTA00000339F.c.02.1	M00001381C:B08	
5096 2/24/98 418 RTA00000410F.a.01.1 M00001631D:B10 73354 5097 3/24/98 458 RTA00000424F.d.12.2 M00001530D:E06 74342 5098 3/24/98 458 RTA00000424F.d.12.2 M00001530D:E06 74342 5098 3/24/98 454 RTA00000424F.d.12.2 M00001530D:E06 74342 5098 3/24/98 454 RTA00000424F.d.12.2 M00001530D:E06 74342 5099 2/24/98 159 RTA00000348F.j.17.1 M00001391D:C06 2641 5100 2/24/98 159 RTA00000346F.m.15.1 M00001632B:E05 3655 5101 2/24/98 162 RTA00000522F.a.08.1 M00001632B:E05 3655 5102 3/24/98 162 RTA0000042F.b.2.1 M00000437A:E04 13767 5104 2/24/98 119 RTA0000042F.b.22.1 M00000437A:E04 13767 5105 3/24/98 12 RTA0000024F.b.22.1 M00001530A:F11 72971 5106 3/24/98 8	5094	2/24/98	1243	RTA00000404F.j.19.1	M00001630D:H10	
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5097 3/24/98 454 RTA00000424F.d.12.2 M00001530D:E06 74342 5098 3/24/98 458 RTA00000424F.d.12.2 M00001530D:E06 74342 5099 2/24/98 159 RTA00000348R.j.17.1 M00001530D:E06 74342 5099 2/24/98 159 RTA00000346F.m.15.1 M00001632B:E05 3655 5100 2/24/98 170 RTA00000422F.n.08.1 M00001632B:E05 38655 5101 3/24/98 162 RTA00000422F.n.08.1 M00001632B:E05 38655 5102 3/24/98 162 RTA00000422F.n.08.1 M00001632B:E05 38655 5103 2/24/98 119 RTA00000422F.n.08.1 M00004037A:E04 13767 5104 2/24/98 119 RTA000000522F.a.23.1 M00001570C:A05 38613 5106 3/24/98 103 RTA00000424F.b.22.1 M00001530A:F11 72971 5106 3/24/98 8 RTA00000441F.g.08.1 M00001530A:F11 72971 5107 2/24/98 3		3/24/98	458	RTA00000424F.d.12.3	M00001530D:E06	
5098 3/24/98 458 RTA00000424F.d.12.3 M00001530D:E06 74342 5098 3/24/98 454 RTA00000424F.d.12.2 M00001530D:E06 74342 5099 2/24/98 159 RTA0000034RB.j.17.1 M00001391D:C06 2641 5100 2/24/98 170 RTA0000034Fm.ls.1 M00004037B:C04 13553 5101 2/24/98 170 RTA0000042Fm.08.1 M00001632B:E05 38655 5102 3/24/98 162 RTA0000042Fm.08.1 M00004037A:E04 13767 5103 2/24/98 119 RTA0000042Fm.08.1 M00004037A:E04 3767 5104 2/24/98 119 RTA0000042Fm.02.1 M0000436D:F02 37472 5105 3/24/98 12 RTA0000042Fm.02.1 M00001530A:F11 72971 5106 3/24/98 8 RTA0000042Fb.22.4 M00001530A:F11 72971 5107 2/24/98 21 RTA0000042Fb.22.4 M00001530A:F11 72971 5108 1/28/98 35 RTA0000044		3/24/98	454	RTA00000424F.d.12.2	M00001530D:E06	
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5100 2/24/98 539 RTA00000346F.m.15.1 M00004037B:C04 13553 5101 2/24/98 162 RTA00000422F.n.08.1 M00001632B:E05 38655 5102 3/24/98 162 RTA00000522F.a.12.1 M00001567A:H05 33515 5104 2/24/98 315 RTA00000419F.p.12.1 M00004037A:E04 13767 5104 2/24/98 119 RTA0000042F.b.25.1 M0000437A:E04 13767 5105 3/24/98 12 RTA00000522F.a.23.1 M00001530A:F11 72971 5106 3/24/98 88 RTA00000424F.b.22.1 M00001530A:F11 72971 5106 3/24/98 21 RTA00000424F.b.22.1 M00003822D:D04 45815 5108 1/28/98 35 RTA00000527F.c.23.1 M00003822C:A07 37742 5110 1/28/98 35 RTA00000527F.c.23.1 M00001392D:H04 0 5111 2/24/98 54 RTA00000399F.o.01.1 M00001535D:C01 38638 5112 2/24/98 63 <td< td=""><td></td><td></td><td>159</td><td>RTA00000348R.j.17.1</td><td>M00001391D:C06</td><td></td></td<>			159	RTA00000348R.j.17.1	M00001391D:C06	
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5102 3/24/98 162 RTA00000522F.a.12.1 M00001567A:H05 33515 5103 2/24/98 315 RTA00000419F.p.12.1 M00004037A:E04 13767 5104 2/24/98 315 RTA00000423F.k.05.1 M000004036D:F02 37472 5105 3/24/98 12 RTA00000522F.a.23.1 M00001570C:A05 38613 5106 3/24/98 88 RTA00000424F.b.22.4 M00001530A:F11 72971 5107 2/24/98 21 RTA0000041F.g.08.1 M00003822D:D04 45815 5108 1/28/98 35 RTA00000527F.c.23.1 M00000382D:D04 45815 5109 3/24/98 39 RTA00000527F.c.23.1 M00000382D:D04 45815 5110 1/28/98 39 RTA00000527F.c.23.1 M00000382D:H04 0 5111 2/24/98 54 RTA00000387F.c.14.3 M00001392D:H04 0 5112 2/24/98 54 RTA0000044F.l.20.2 M00001639E.C01 3484 5113 1/28/98 82 RTA00				RTA00000422F.n.08.1	M00001632B:E05	38655
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5106 3/24/98 88 RTA00000424F.b.22.4 M00001530A:F11 72971 5107 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 5108 1/28/98 35 RTA00000191AF.n.17.1 M000004091B:D11 7848 5109 3/24/98 39 RTA00000527F.c.23.1 M00003822C:A07 37742 5110 1/28/98 43 RTA00000179AF.c.14.3 M00001392D:H04 0 5111 2/24/98 54 RTA00000399F.o.01.1 M00001595C:E01 3055 5112 2/24/98 63 RTA0000044F.l.20.2 M00001639B:H05 38638 5113 1/28/98 82 RTA00000183AF.l.18.1 M00001595C:E01 3484 5114 3/24/98 84 RTA00000527F.k.18.1 M00001595D:C01 3484 5115 1/28/98 99 RTA00000184AF.d.8.1 M00001548A:A08 4393 5116 2/24/98 10 RTA00000034F.n.19.1 M00001473C:D09 7801 5117 2/24/98 104 RTA0			103	RTA00000424F.b.22.1	M00001530A:F11	72971
5107 2/24/98 21 RTA00000411F.g.08.1 M00003822D:D04 45815 5108 1/28/98 35 RTA00000191AF.n.17.1 M00004091B:D11 7848 5109 3/24/98 39 RTA00000527F.c.23.1 M00003822C:A07 37742 5110 1/28/98 43 RTA00000179AF.c.14.3 M00001392D:H04 0 5111 2/24/98 54 RTA00000399F.o.01.1 M00001595C:E01 3055 5112 2/24/98 63 RTA0000044F.l.20.2 M00001639B:H05 38638 5113 1/28/98 82 RTA00000183AF.l.18.1 M00001535D:C01 3484 5114 3/24/98 84 RTA00000527F.k.18.1 M00001535D:C01 3484 5115 1/28/98 99 RTA00000184AF.d.8.1 M00001548A:A08 4393 5116 2/24/98 99 RTA00000339F.o.23.1 M00001473C:D09 7801 5118 2/24/98 104 RTA0000034F.n.03.1 M00001473C:D09 7801 5120 2/24/98 105 RTA000			88	RTA00000424F.b.22.4	M00001530A:F11	72971
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5116 2/24/98 99 RTA00000420F.m.19.1 M00005254D:B08 0 5117 2/24/98 100 RTA00000339F.o.23.1 M00001473C:D09 7801 5118 2/24/98 104 RTA00000421F.n.03.1 M00001675C:A04 1638 5119 2/24/98 105 RTA00000346F.d.08.1 M00001671A:A10 39955 5120 2/24/98 114 RTA00000341F.m.21.1 M00004051D:E01 0 5121 1/28/98 137 RTA00000181AF.m.4.3 M00001455A:E09 13238 5122 1/28/98 162 RTA00000201F.e.15.1 M00004444B:D11 9960 5123 1/28/98 170 RTA00000197AF.d.23.1 M00001453A:E11 16130 5124 1/28/98 206 RTA00000181AF.o.04.2 M00001457C:C12 22205 5125 1/28/98 209 RTA00000182AF.c.5.1 M00001539D:E10 5790 5127 2/24/98 215 RTA0000043F.j.18.1 M00001615C:F03 0 5129 1/28/98 230 RT	5114	3/24/98	84			
5117 2/24/98 100 RTA00000339F.o.23.1 M00001473C:D09 7801 5118 2/24/98 104 RTA00000421F.n.03.1 M00001675C:A04 1638 5119 2/24/98 105 RTA00000346F.d.08.1 M00001671A:A10 39955 5120 2/24/98 114 RTA00000341F.m.21.1 M00004051D:E01 0 5121 1/28/98 137 RTA00000181AF.m.4.3 M00001455A:E09 13238 5122 1/28/98 162 RTA00000201F.e.15.1 M00004444B:D11 9960 5123 1/28/98 170 RTA00000197AF.d.23.1 M00001453A:E11 16130 5124 1/28/98 206 RTA00000181AF.o.04.2 M00001457C:C12 22205 5125 1/28/98 209 RTA00000182AF.c.5.1 M00001464D:F06 6397 5126 2/24/98 215 RTA0000043F.j.18.1 M00001539D:E10 5790 5127 2/24/98 219 RTA00000185AR.b.18.1 M00001575B:C09 12171 5130 3/24/98 245	5115	1/28/98	99			
5118 2/24/98 104 RTA00000421F.n.03.1 M00001675C:A04 1638 5119 2/24/98 105 RTA00000346F.d.08.1 M00001671A:A10 39955 5120 2/24/98 114 RTA00000341F.m.21.1 M00004051D:E01 0 5121 1/28/98 137 RTA00000181AF.m.4.3 M00001455A:E09 13238 5122 1/28/98 162 RTA00000201F.e.15.1 M00004444B:D11 9960 5123 1/28/98 170 RTA00000197AF.d.23.1 M00001453A:E11 16130 5124 1/28/98 206 RTA00000181AF.o.04.2 M00001457C:C12 22205 5125 1/28/98 209 RTA00000182AF.c.5.1 M00001464D:F06 6397 5126 2/24/98 215 RTA0000043F.j.18.1 M00001539D:E10 5790 5127 2/24/98 219 RTA00000185AR.b.18.1 M0000167C:F03 0 5129 1/28/98 230 RTA00000185AR.b.18.1 M00001670A:F09 75204 5130 3/24/98 245	5116	· 2/24/98	99			
5119 2/24/98 105 RTA00000346F.d.08.1 M00001671A:A10 39955 5120 2/24/98 114 RTA00000341F.m.21.1 M00004051D:E01 0 5121 1/28/98 137 RTA00000181AF.m.4.3 M00001455A:E09 13238 5122 1/28/98 162 RTA00000201F.e.15.1 M00004444B:D11 9960 5123 1/28/98 170 RTA00000197AF.d.23.1 M00001453A:E11 16130 5124 1/28/98 206 RTA00000181AF.o.04.2 M00001457C:C12 22205 5125 1/28/98 209 RTA00000182AF.c.5.1 M00001464D:F06 6397 5126 2/24/98 215 RTA0000043F.j.18.1 M00001539D:E10 5790 5127 2/24/98 219 RTA00000198AF.g.3.1 M00001615C:F03 0 5129 1/28/98 230 RTA00000185AR.b.18.1 M00001575B:C09 12171 5130 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204	5117	2/24/98	100			
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5121 1/28/98 137 RTA00000181AF.m.4.3 M00001455A:E09 13238 5122 1/28/98 162 RTA00000201F.e.15.1 M00004444B:D11 9960 5123 1/28/98 170 RTA00000197AF.d.23.1 M00001453A:E11 16130 5124 1/28/98 206 RTA00000181AF.o.04.2 M00001457C:C12 22205 5125 1/28/98 209 RTA00000182AF.c.5.1 M00001464D:F06 6397 5126 2/24/98 215 RTA0000043F.j.18.1 M00001539D:E10 5790 5127 2/24/98 219 RTA00000419F.c.18.1 M00003819D:B11 41394 5128 1/28/98 229 RTA00000185AR.b.18.1 M0000167C:F03 0 5129 1/28/98 230 RTA00000185AR.b.18.1 M00001670A:F09 75204 5130 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204	5119	2/24/98	105			
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5124 1/28/98 209 RTA00000182AF.c.5.1 M00001464D:F06 6397 5126 2/24/98 215 RTA00000403F.j.18.1 M00001539D:E10 5790 5127 2/24/98 219 RTA00000419F.c.18.1 M00003819D:B11 41394 5128 1/28/98 229 RTA00000198AF.g.3.1 M00001615C:F03 0 5129 1/28/98 230 RTA00000185AR.b.18.1 M00001575B:C09 12171 5130 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204	5123					
5126 2/24/98 215 RTA00000403F.j.18.1 M00001539D:E10 5790 5127 2/24/98 219 RTA00000419F.c.18.1 M00003819D:B11 41394 5128 1/28/98 229 RTA00000198AF.g.3.1 M00001615C:F03 0 5129 1/28/98 230 RTA00000185AR.b.18.1 M00001575B:C09 12171 5130 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204	5124					
5127 2/24/98 219 RTA00000419F.c.18.1 M00003819D:B11 41394 5128 1/28/98 229 RTA00000198AF.g.3.1 M00001615C:F03 0 5129 1/28/98 230 RTA00000185AR.b.18.1 M00001575B:C09 12171 5130 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204						
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5129 1/28/98 230 RTA00000185AR.b.18.1 M00001575B:C09 12171 5130 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204						
5130 3/24/98 245 RTA00000522F.p.09.1 M00001670A:F09 75204						
5150 5/2 1/70 210 2001						
5131 2/24/98 258 RTA00000406F.k.15.1 M0000390/C:C04 38549				-		
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	Priority	Priority			
	Appln	Appln			0551
5132	1/28/98	262	RTA00000186AF.c.17.1	M00001619D:G05	8551
5133	1/28/98	269	RTA00000183AF.k.13.1	M00001534B:C12	0
5134	1/28/98	276	RTA00000198AF.j.15.1	M00001653B:E09	4369
5135	2/24/98	281	RTA00000411F.l.13.1	M00003857C:C09	43114
5136	1/28/98	284	RTA00000199F.d.10.2	M00003808C:B05	22049
5137	1/28/98	292	RTA00000199AF.m.18.1	M00003939C:F04	0
5138	1/28/98	297	RTA00000178AF.f.9.3	M00001371C:E09	7172
5139	2/24/98	301	RTA00000401F.m.23.1	M00003914C:C02	2801
5140	1/28/98	302	RTA00000186AF.d.1.2	M00001621C:C08	40044
5141	1/28/98	315	RTA00000199R.d.23.1	M00003815D:H09	37477
5142	2/24/98	315	RTA00000419F.p.12.1	M00004037A:E04	13767
5143	1/28/98	321	RTA00000181AR.b.21.1	M00001444C:D05	3266
5144	3/24/98	323	RTA00000524F.c.12.1	M00005218B:D09	0
5145	1/28/98	329	RTA00000186AF.b.9.1	M00001616C:F07	0
5146	1/28/98	334	RTA00000181AR.b.21.3	M00001444C:D05	3266
5147	2/24/98	334	RTA00000408F.p.05.1	M00001575B:B02	9649
5148	1/28/98	335	RTA00000182AF.e.3.2	M00001468B:H06	0
5149	1/28/98	336	RTA00000186AF.f.24.1	M00001629B:E06	0
5150	2/24/98	341	RTA00000412F.g.20.2	M00003972C:F08	25018
5151	2/24/98	343	RTA00000422F.g.21.1	M00001583A:F07	17232
5152	1/28/98	347	RTA00000199F.b.03.2	M00003779B:E12	38340
5153	2/24/98	354	RTA00000404F.c.03.2	M00001592C:F11	39198
5154	1/28/98	361	RTA00000177AR.g.16.4	M00001347A:B10	13576
5155	1/28/98	364	RTA00000187AF.g.13.1	M00001676C:C11	2991
5156	2/24/98	377	RTA00000346F.i.01.1	M00003797A:D06	22260
5157	2/24/98	389	RTA00000411F.c.02.1	M00001677B:B04	72852
5158	2/24/98	403	RTA00000403F.d.22.1	M00001473A:A07	10692
5159	1/28/98	407	RTA00000178AF.e.20.1	M00001370D:E12	3135
5160	1/28/98	422	RTA00000189AF.b.12.1	M00003829B:G03	17233
5161	2/24/98	429	RTA00000422F.c.17.1	M00004099D:F01	1360
5162	2/24/98	431	RTA00000399F.j.15.1	M00001578C:G06	1261
5163	1/28/98	439	RTA00000185AF.d.14.2	M00001579D:G07	8071
5164	2/24/98	448	RTA00000127A.a.3.1	M00001552A:H10	13232
5165	2/24/98	450	RTA00000118A.a.23.1	M00001395A:H02	3500
5166	1/28/98	451	RTA00000200AF.b.20.1	M00004043A:D02	40403
5167	2/24/98	455	RTA00000399F.d.23.1	M00001481B:A07	3310
5168	1/28/98	475	RTA00000187AR.m.3.3	M00001682C:B12	17055
5169	3/24/98	475	RTA00000427F.i.06.1	M00004097B:D03	41450
5170	3/24/98	477	RTA00000527F.l.21.1	M00003983D:H02	36439
5171	1/28/98	480	RTA00000181AF.o.08.2	M00001457C:H12	849
5172	3/24/98	480	RTA00000424F.d.17.3	M00001455A:E11	73958
5173	3/24/98	481	RTA00000523F.j.02.1	M00003857A:H10	62853
5174	1/28/98	483	RTA00000192AF.h.19.1	M00004162C:A07	4642
5175	2/24/98	489	RTA00000406F.j.19.1	M00003906A:F12	1685
5176	1/28/98	501	RTA00000200R.k.11.1	M00004197C:F03	9796
5177	2/24/98	502	RTA00000341F.d.08.1	M00003824C:D07	0
5178	2/24/98	508	RTA00000420F.i.20.1	M00005101C:E12	0
5179	1/28/98	510	RTA00000178AF.n.23.1	M00001387B:E02	3298
5180	1/28/98	511	RTA00000196AF.g.10.1	M00001376B:A02	12498
5181	2/24/98	519	RTA00000404F.I.10.1	M00001638B:F10	23136

Applin	SEQ ID NO:	Filing Date of Priority	SEQ ID NO: in Priority	Sequence Name	Clone Name	Cluster ID
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5183 1/28/98 525 RTA00000198AF.c.7.1 M00001575D:GG5 19181 5184 1/28/98 526 RTA00000198AF.c.20.1 M00001585A:D06 5865 5185 1/28/98 527 RTA00000198AE.c.21. M00001684B:G03 38469 5186 1/28/98 529 RTA00000185AF.d.24.2 M00001582D:F05 0 5187 1/28/98 540 RTA00000179AF.b.10.3 M00001591D:D10 0 5189 1/28/98 541 RTA00000197AR.b.16.1 M00001445C:A08 0 5190 1/28/98 547 RTA00000197AR.b.16.1 M0000138B:E02 3575 5191 2/24/98 547 RTA0000018AF.g.14.1 M0000137B:D:G03 35258 5192 1/28/98 548 RTA00000133A-d.22.1 M00001497B:D:G03 35258 5193 1/28/98 550 RTA0000013AF.g.14.1 M00001499A:A05 3728 5194 2/24/98 569 RTA0000019AF.g.12.3.1 M00001499A:G11 11797 5195 1/28/98 570	5182			RTA00000419F f 23 1	M00003840D:H10	65002
5184 1/28/98 526 RTA00000185AF.e.20.1 M00001685A:D06 5865 5185 1/28/98 527 RTA00000178AF.b.13.1 M00001684B:G03 38469 5186 1/28/98 529 RTA00000178AF.b.10.3 M00001354A:E11 3114 5187 1/28/98 540 RTA00000179AF.b.10.3 M00001391D:D10 0 5189 1/28/98 541 RTA00000197AR.b.16.1 M0000138B:D202 3575 5190 1/28/98 543 RTA0000019FAB.b.10.1 M00001345D:C08 395 5191 2/24/98 547 RTA0000019FAB.b.02.1 M0000345D:G08 3985 5192 1/28/98 548 RTA0000019FAB.c.23.3 M0000139D:G03 35258 5193 1/28/98 550 RTA0000019FAB.c.23.3 M0000139D:G03 35258 5194 2/24/98 555 RTA0000019AB.c.23.1 M00001412A:E04 11797 5195 1/28/98 569 RTA0000019AB.c.23.1 M00001412A:E04 11797 5196 1/28/98 570						
5185 1/28/98 527 RTA00000198R.m.23.1 M00001684B:G03 38469 5186 1/28/98 529 RTA000001158AF.b.13.1 M00001684B:G03 38469 5187 1/28/98 530 RTA000001158AF.d.24.2 M000001582D:F05 0 5188 1/28/98 540 RTA00000179AF.b.10.3 M00001391D:D10 0 5189 1/28/98 541 RTA00000197AR.b.16.1 M00001338B:E02 3575 5190 1/28/98 544 RTA00000197AF.b.2.1 M00001338B:E02 3575 5191 1/28/98 547 RTA00000183AF.g.14.1 M00001338B:E02 3575 5192 1/28/98 550 RTA00000183AF.g.14.1 M00001513D:A03 0 5194 1/28/98 550 RTA00000183AF.g.1.1 M00001499A:A05 3788 5195 1/28/98 570 RTA00000192AF.f.3.1 M00001499A:A05 3788 5197 1/28/98 576 RTA0000019AF.g.1.1 M00001499A:A05 3788 5198 1/28/98 576						5865
5186 1/28/98 529 RTA00000178AF.b.13.1 M00001364A:E11 3114 5187 1/28/98 530 RTA00000179AF.b.10.3 M000013891P:D10 0 5188 1/28/98 540 RTA00000179AF.b.10.3 M00001391P:D10 0 5189 1/28/98 541 RTA0000019FAB.b.16.1 M00001345C:A08 0 5190 1/28/98 547 RTA00000119Fh.02.1 M0000138E5D:G08 3575 5191 2/24/98 547 RTA00000119Fh.02.1 M0000139TB:G03 35258 5192 1/28/98 550 RTA00000119AF.f.23.3 M0000139TB:G03 35258 5193 1/28/98 550 RTA0000019AF.c.23.1 M00001499A:G11 11797 5195 1/28/98 569 RTA0000019AF.c.23.1 M00001499A:G11 11797 5196 1/28/98 570 RTA0000013AF.a.19.2 M00001499A:A05 3788 5197 1/28/98 575 RTA00000149AF.a.21 M0000149A:A05 378 5198 1/28/98 575						
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5214 1/28/98 704 RTA00000190AF.o.12.1 M00003972D:C09 3438 5215 1/28/98 723 RTA00000183AF.p.24.1 M00001543C:F01 3116 5216 2/24/98 733 RTA00000405F.d.18.1 M00001662C:B02 10494 5217 1/28/98 739 RTA00000181AF.p.12.3 M00001460C:H02 22204 5218 1/28/98 742 RTA00000177AF.m.1.1 M00001353D:D10 14929 5219 2/24/98 774 RTA00000403F.e.24.1 M00001476B:D10 16432 5220 2/24/98 775 RTA00000405F.c.22.1 M00001660C:B06 39053 5221 2/24/98 790 RTA00000345F.n.08.1 M00001517A:B11 0 5222 2/24/98 816 RTA00000354R.n.04.1 M00003808C:B05 22049 5223 2/24/98 851 RTA0000043F.d.07.1 M00001678B:B12 0 5224 2/24/98 871 RTA0000043F.d.23.1 M00001678B:B12 74567 5227 2/24/98 914						-
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5218 1/28/98 742 RTA00000177AF.m.1.1 M00001353D:D10 14929 5219 2/24/98 774 RTA00000403F.e.24.1 M00001476B:D10 16432 5220 2/24/98 775 RTA00000405F.c.22.1 M00001660C:B06 39053 5221 2/24/98 790 RTA00000345F.n.08.1 M00001517A:B11 0 5222 2/24/98 816 RTA00000354R.n.04.1 M00003808C:B05 22049 5223 2/24/98 829 RTA0000041F.m.11.1 M00001678B:B12 0 5224 2/24/98 851 RTA00000423F.d.07.1 M00001678B:B12 0 5225 2/24/98 871 RTA0000043F.f.23.1 M00001674P:E01 39223 5226 2/24/98 877 RTA00000138A.m.15.1 M00001624A:A03 41603 5228 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 5229 2/24/98 924 RTA00000354R.m.02.1 M00003890B:C08 12766 5230 2/24/98 940						
5219 2/24/98 774 RTA00000403F.e.24.1 M00001476B:D10 16432 5220 2/24/98 775 RTA00000405F.c.22.1 M00001660C:B06 39053 5221 2/24/98 790 RTA00000345F.n.08.1 M00001517A:B11 0 5222 2/24/98 816 RTA00000354R.n.04.1 M00003808C:B05 22049 5223 2/24/98 829 RTA00000411F.m.11.1 M00003867A:D12 73196 5224 2/24/98 851 RTA00000423F.d.07.1 M00001678B:B12 0 5225 2/24/98 871 RTA0000043F.f.23.1 M00001479C:E01 39223 5226 2/24/98 877 RTA00000418F.m.22.1 M00001654D:E12 74567 5227 2/24/98 914 RTA00000138A.m.15.1 M00001548A:G01 79474 5229 2/24/98 924 RTA00000354R.m.02.1 M00003890B:C08 12766 5230 2/24/98 940 RTA00000414F.f.17.1 M00005260A:F04 0						
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5224 2/24/98 851 RTA00000423F.d.07.1 M00001678B:B12 0 5225 2/24/98 871 RTA00000403F.f.23.1 M00001479C:E01 39223 5226 2/24/98 877 RTA00000418F.m.22.1 M00001654D:E12 74567 5227 2/24/98 914 RTA00000138A.m.15.1 M00001624A:A03 41603 5228 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 5229 2/24/98 924 RTA00000354R.m.02.1 M00003890B:C08 12766 5230 2/24/98 940 RTA00000414F.f.17.1 M00005260A:F04 0						
5225 2/24/98 871 RTA00000403F.f.23.1 M00001479C:E01 39223 5226 2/24/98 877 RTA00000418F.m.22.1 M00001654D:E12 74567 5227 2/24/98 914 RTA00000138A.m.15.1 M00001624A:A03 41603 5228 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 5229 2/24/98 924 RTA00000354R.m.02.1 M00003890B:C08 12766 5230 2/24/98 940 RTA00000414F.f.17.1 M00005260A:F04 0						
5226 2/24/98 877 RTA00000418F.m.22.1 M00001654D:E12 74567 5227 2/24/98 914 RTA00000138A.m.15.1 M00001624A:A03 41603 5228 2/24/98 923 RTA00000126A.d.19.1 M00001548A:G01 79474 5229 2/24/98 924 RTA00000354R.m.02.1 M00003890B:C08 12766 5230 2/24/98 940 RTA00000414F.f.17.1 M00005260A:F04 0						
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5229 2/24/98 924 RTA00000354R.m.02.1 M00003890B:C08 12766 5230 2/24/98 940 RTA00000414F.f.17.1 M00005260A:F04 0						
5230 2/24/98 940 RTA00000414F.f.17.1 M00005260A:F04 0						

SEQ ID	Filing	SEQ ID	Sequence Name	Clone Name	Cluster
NO:	Date of	NO: in			ID
	Priority	Priority			
	Appln	Appln			
5232	2/24/98	1013	RTA00000404F.b.18.1	M00001592A:H05	13669
5233	2/24/98	1037	RTA00000339F.I.12.1	M00001450A:G11	7711
5234	2/24/98	1055	RTA00000346F.a.04.1	M00001607B:C05	5382
5235	2/24/98	1070	RTA00000346F.n.22.1	M00004137A:D06	0
5236	2/24/98	1096	RTA00000421F.a.06.1	M00001589C:A11	2385
5237	2/24/98	1125	RTA00000118A.n.5.1	M00001451A:C10	0
5238	2/24/98	1128	RTA00000423F.a.02.3	M00001656B:A08	39210
5239	2/24/98	1129	RTA00000401F.m.07.1	M00003907D:F11	2893
5240	2/24/98	1136	RTA00000406F.c.05.1	M00003870A:H01	22077
5241	2/24/98	1142	RTA00000418F.i.06.1	M00001591B:B06	75151
5242	2/24/98	1145	RTA00000423F.k.21.2	M00003984D:B08	37499
5243	2/24/98	1149	RTA00000339F.b.02.1	M00001344B:F12	0
5244	2/24/98	1166	RTA00000347F.h.01.1	M00004040A:G12	12043
5245	2/24/98	1177	RTA00000126A.b.9.1	M00001547A:F11	81279
5246	2/24/98	1187	RTA00000120A.c.19.1	M00001464A:B03	81016
5247	2/24/98	1203	RTA00000413F.e.10.1	M00004092C:B03	31033
5248	2/24/98	1205	RTA00000419F.k.05.1	M00003871C:E04	11757
5249	2/24/98	1230	RTA00000399F.j.14.1	M00001578C:F05	16942
5250	2/24/98	1233	RTA00000418F.I.02.1	M00001641C:C05	39316
5251	2/24/98	1248	RTA00000419F.o.07.1	M00003986C:E09	14059
5252	2/24/98	1261	RTA00000404F.m.17.2	M00001643B:E05	0

Table 2

Table	Table 2								
		learest Neighbor		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)					
	,	astN vs. Genbank)							
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE			
2503	AB011149	Homo sapiens mRNA for KIAA0577 protein, complete	0	3043678	(AB011149) KIAA0577 protein [Homo sapiens]	1e-096			
2504	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>			
2505	Z59973	H.sapiens CpG DNA, clone 184b10, forward read cpg184b10.ft1a.	1e-009	<none></none>	<none></none>	<none></none>			
2506	AJ000742	Homo Sapiens hisH1 gene, 5' UTR	2e-016	<none></none>	<none></none>	<none></none>			
2507	U10324	Human nuclear factor NF90 mRNA, complete cds.	3e-009	1729881	TETRACYCLINE RESISTANCE PROTEIN, CLASS H (TETA(H)) >gi 392873 (U00792) tetracycline resistance protein [Pasteurella multocida]	9.3			
2508	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	1890128	(U89949) folate binding protein [Sus scrofa]	7.3			
2509	M15657	Human aldolase B (ALDOB) gene, exons 2 through 6.	0.002	<none></none>	<none></none>	<none></none>			
2510	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-008	<none></none>	<none></none>	<none></none>			
2511	U39722	Mycoplasma genitalium section 44 of 51 of the complete genome	0.043	2773162	(AF039595) sulfonylurea receptor 1B [Rattus norvegicus]	10			
2512	AB012174	Homo sapiens DNA, anonymous heat-stable fragment RP7-1B	7e-017	<none></none>	<none></none>	<none></none>			

•	<u> </u>	learest Neighbor		Nearest Neighbor			
	(BI	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID						370375	
2513	AB012174	Homo sapiens	7e-017	<none></none>	<none></none>	<none></none>	
		DNA, anonymous					
		heat-stable					
2514	U95094	fragment RP7-1B Xenopus laevis	7e-007	2984585	(AC004472)	1e-013	
2314	093094	XL-INCENP	70-007	2901303	P1.11659 4		
		(XL-INCENP)			[Homo sapiens]		
		mRNA, complete					
		cds					
2515	AF061016	Homo sapiens	0	3127127	(AF061016) UDP-	7e-035	
		UDP-glucose			glucose		
		dehydrogenase			dehydrogenase		
		(UGDH) mRNA,			[Homo sapiens]		
		complete cds			dehydrogenase [Homo sapiens]		
2516	U95102	Xenopus laevis	6e-005	2983872	(AE000742)	1.5	
2316	093102	mitotic	06-003	2903072	putative protein	1.5	
		phosphoprotein			[Aquifex aeolicus]		
		90 mRNA,					
		complete cds					
2517	X13293	Human mRNA	3e-019	127584	MYB-RELATED	0.0002	
		for B-myb gene			PROTEIN B (B-		
					MYB) human >gi 29472		
					(X13293) B-myb		
					protein (AA 1-		
-					700) [Homo		
					sapiens]		
2518	Y10183	H.sapiens mRNA	0	3882036	(AJ010405)	2.5	
		for MEMD			hypothetical		
	160000	protein	4 000	2051770	protein	7.0	
2519	M90297	Human	4e-023	2851668	HYPOTHETICAL OUTER	7.8	
		glucokinase (GCK) gene,			MEMBRANE		
		exon 1 and 5'			USHER		
		flanking region.			PROTEIN IN		
					RIBB-GLGS		
					INTERGENIC		
					REGION		
				161	PRECURSOR	210272	
2520	V00436	Gallus gallus	4.4	<none></none>	<none></none>	<none></none>	
		fragment of gene X of ovalbumin					
		family coding for					
		the first leader					
		exon.					
L	L	<u> </u>			*		

		Nearest Neighbor		Nearest Neighbor			
	(Bl	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID			6 006	2000011	(A E070051)	(0	
2521	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	3800811	(AF072251) methyl-CpG binding protein 2 [Mus musculus]	6.9	
2522	Y09540	H.sapiens AHSG gene, partial	2e-007	2135357	HLA class I alpha chain - human (fragment) sapiens]	3.1	
2523	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	3e-007	<none></none>	<none></none>	<none></none>	
2524	D87438	Human mRNA for KIAA0251 gene, partial cds	1e-011	<none></none>	<none></none>	<none></none>	
2525	AE001203	Treponema pallidum section 19 of 87 of the complete genome	0.42	<none></none>	<none></none>	<none></none>	
2526	U47322	Cloning vector DNA, complete sequence.	2e-036	987050	(X65335) lacZ gene product [unidentified cloning vector]	4e-008	
2527	M97287	Human MAR/SAR DNA binding protein (SATB1) mRNA, complete cds. > :: gb I58691 I58691 Sequence 1 from patent US 5652340	0	417747	DNA-BINDING PROTEIN SATB1 (SPECIAL AT- RICH SEQUENCE BINDING PROTEIN 1) protein SATB1 - human >gi 337811 (M97287) putative [Homo sapiens]	2e-009	
2528	AF005355	Oryctolagus cuniculus translation initiation factor eIF2C mRNA, complete cds	1e-094	3253159	(AF005355) translation initiation factor eIF2C	2e-084	
2529	L16978	Anadara trapezia beta globin gene, complete cds.	0.11	<none></none>	<none></none>	<none></none>	
2530	M24191	Human beta globulin pseudogene, clone 46B	0.013	3878519	(Z92806) K10G4.7 [Caenorhabditis elegans]	0.6	

	N	learest Neighbor		Nearest Neighbor			
	(BlastN vs. Genbank)			(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2531	AF047611	Euroglyphus maynei group 1 allergen Eur m 1 0102	0.12	<none></none>	<none></none>	<none></none>	
2532	AE001372	Plasmodium falciparum chromosome 2, section 9 of 73 of the complete sequence	0.002	<none></none>	<none></none>	<none></none>	
2533	J04700	Homo sapiens calcium- dependent protease large subunit (CANPmL) gene, promoter region and exon 1.	0.014	<none></none>	<none></none>	<none></none>	
2534	AF038958	Homo sapiens synaptic glycoprotein SC2 spliced variant mRNA, complete cds	4e-086	2144098	SC2 - rat >gi 256994 bbs 11 5268 (S45663) SC2=synaptic glycoprotein [rats, brain, Peptide, 308 aa]	1e-033	
2535	L13434	Human chromosome 3p21.1 gene sequence, complete cds.	8e-008	1085432	mucin (clone PGM-2A) - pig	4.3	
2536	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	3873713	(Z74026) cDNA EST yk452h4.3 comes from this gene; cDNA EST yk452h4.5 comes from this gene	4e-010	
2537	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	<none></none>	<none></none>	<none></none>	
2538	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	386644	type Ia hair keratin a3 [human, Peptide, 404 aa] >gi 3724101 gnl PI D e1330425 (Y16788) keratin, type I [Homo sapiens]	1.9	

	<u>N</u>	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2539	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0005	<none></none>	<none></none>	<none></none>	
2540	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
2541	U79248	Human clone 23826 mRNA sequence	6e-005	<none></none>	<none></none>	<none></none>	
2542	D44598	Saccharomyces cerevisiae chromosome VI phage 4121	1e-010	2828280	(AL021687) putative protein [Arabidopsis thaliana] >gi 2832633 gnl PI D e1249651 (AL021711) putative protein [Arabidopsis thaliana]	6e-060	
2543	X64037	H.sapiens mRNA for RNA polymerase II associated protein RAP74	0	35871	(X64002) RAP74 [Homo sapiens] >gi 228483 prf 18 04353A transcription factor RAP74 [Homo sapiens]	4e-049	
2544	M18857	A.californica nuclear polyhedrosis virus ORFs encoding a delayed early protein and two late protein, complete cds.	0.38	3169096	(AL023706) hypothetical protein	3e-029	
2545	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-006	<none></none>	<none></none>	<none></none>	
2546	L22403	Homo sapiens DNA sequence, repeat region.	1e-020	<none></none>	<none></none>	<none></none>	
2547	L22403	Homo sapiens DNA sequence, repeat region.	1e-020	<none></none>	<none></none>	<none></none>	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2548	D38417	Mouse mRNA for arylhydrocarbon receptor, complete cds	3e-028	<none></none>	<none></none>	<none></none>	
2549	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
2550	X04754	Drosophila yolk polypeptide gene YP3	1e-012	2500649	PROBABLE RNA 3'-TERMINAL PHOSPHATE CYCLASE (RNA- 3'-PHOSPHATE CYCLASE) (RNA- CYCLASE)	1e-022	
2551	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-011	<none></none>	<none></none>	<none></none>	
2552	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
2553	U49169	Dictyostelium discoideum V- ATPase A subunit (vatA) mRNA, complete cds	0.13	586429	VERY HYPOTHETICAL 13.2 KD PROTEIN IN PTC3-SAS3 INTERGENIC REGION >gi 626813 pir S4 5788 probable membrane protein YBL053w - yeast (Saccharomyces cerevisiae) >gi 536079 (Z35814) ORF YBL053w	1.1	
2554	M22462	Chicken protein p54 (ets-1) mRNA, complete cds.	1.1	2078531	(U89506) Mlark [Mus musculus]	5.6	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2555	U73664	Human t(11;14)(q13;q32) breakpoint junction sequence	0.37	2909381	(Y16569) ORF [Mycobacterium tuberculosis]	3.3	
2556	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	3025166	PUTATIVE NUCLEOSIDE TRANSPORTER YEGT >gi 1736823 gnl PI D d1016692 (D90848) Nucleoside permease NupG (Nucleoside- transport system protein NupG). [Escherichia coli] >gi 1788415 (AE000299) putative nucleoside permease protein [Escherichia coli]	1.4	
2557	U09210	Human vesicular acetylcholine transporter mRNA, complete cds.		3176395	(AB015041) PIF1 [Caenorhabditis elegans]	1e-006	
2558	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.002	540271	(U14635) similar to GABA and glycine receptors	1e-020	
2559		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
2560	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	<none></none>	<none></none>	<none></none>	
2561	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	1788739	(AE000327) orf, hypothetical protein [Escherichia coli]	6.8	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)	•	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2562	AF073710	Homo sapiens regulator of G-protein signaling 9 mRNA, complete cds	1e-013	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.38	
2563	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.1	
2564	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-007	<none></none>	<none></none>	<none></none>	
2565	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-015	<none></none>	<none></none>	<none></none>	
2566	M98502	Mus musculus protein encoding twelve zinc finger proteins (pMLZ-4) mRNA, complete cds.	2e-017	2370153	(Y13374) putative prenylated protein prenylated protein Homo sapiens] >gi 3360403 (AF052096) putative prenylated protein Homo sapiens]	7.3	
2567	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.5	
2568	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-008	2580433	(D76414) ppGpp hydrolase [Staphylococcus aureus]	2.4	
2569	X82206	H.sapiens mRNA for alpha- centractin	4e-085	2909479	(AL021930) hypothetical protein Rv0290	1.4	
2570	Z68758	Human DNA sequence from cosmid cN85E10 on chromosome 22q11.2-qter	8e-009	1082778	secretory phospholipase A2 receptor precursor, transmembrane form - human >gi 862375	7.1	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2571	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0005	2583019	(AF022724) ARIX homeodomain protein [Homo sapiens]	0.64	
2572	L19637	Arabidopsis thaliana adenine phosphoribosyltra nsferase (apt) gene, complete cds.	0.12	<none></none>	<none></none>	<none></none>	
2573	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-008	388057	(L22982) merozoite surface protein-1 [Plasmodium chabaudi]	6.9	
2574	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0005	3913436	PROBABLE ATP- DEPENDENT RNA HELICASE A (NUCLEAR DNA HELICASE II) (NDH II)	9.5	
2575	AJ005698	Homo sapiens mRNA for poly(A)-specific ribonuclease	3e-011	3776076	(AJ005698) poly(A)-specific ribonuclease [Homo sapiens]	0.28	
2576	Z96602	H.sapiens telomeric DNA sequence, clone 3QTEL015, read 3QTELOO015.se	2e-006	2407641	(AF018956) neuropilin [Homo sapiens]	1.4	
2577	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	3880672	(AL032633) cDNA EST EMBL:T00127 comes from this gene; cDNA EST EMBL:T01189 comes from this gene [Caenorhabditis elegans]	0.82	
2578	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	2088843	(AF003386) F59E12.9 gene product [Caenorhabditis elegans]	3.5	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2579	U95315	Mycobacterium gordonae IS1511 transposase and Tn554 tpna transposase homolog genes, complete cds	3.8	<none></none>	<none></none>	<none></none>	
2580	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	9.6	
2581	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-013	<none></none>	<none></none>	<none></none>	
2582	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-008	<none></none>	<none></none>	<none></none>	
2583	U85193	Human nuclear factor I-B2 (NFIB2) mRNA, complete cds	2e-038	<none></none>	<none></none>	<none></none>	
2584	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.2	
2585	U67532	Methanococcus jannaschii section 74 of 150 of the complete genome	0.005	1938410	(U97000) No definition line found [Caenorhabditis elegans]	4.5	
2586	X65319	Cloning vector pCAT-Enhancer	3e-081	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	
2587	AB006534	Homo sapiens mRNA for hepatocyte growth factor activator inhibitor type 2, complete cds	e-103	2065529	(U78095) bikunin [Homo sapiens]	3e-025	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)			s. Non-Redundant Pro		
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID			6 006	3152559	(AC002986)	6e-008	
2588	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006		Similarity to A. thaliana gene product F21M12.20, gb AC000132. EST gb Z25651 comes from this gene. [Arabidopsis thaliana]	3e-017	
2589	X82829	B.taurus mRNA for nuclear DNA helicase II	9e-009	1353239	(U10245) putative RNA helicase A [Arabidopsis thaliana]		
2590	AE001366	Plasmodium falciparum chromosome 2, section 3 of 73 of the complete sequence	0.047	<none></none>	<none></none>	<none></none>	
2591	D78572	House mouse; Musculus domesticus mRNA for membrane glycoprotein, complete cds > :: dbj E12950 E129 50 cDNA GA3- 43 encoding novel polypeptide which appear when differentiate from embryo-tumor cell P19 to nerve cell		1545807	(D78572) membrane glycoprotein [Mus musculus]	1e-026	
2592	M77130	H.sapiens (clone B7) hY4 Ro RNA pseudogene.	4e-011	629174	cellulose 1,4-beta- cellobiosidase (EC 3.2.1.91) - Clostridium thermocellum >gi 530014 (X80993) cellulose 1,4-beta- cellobiosidase [Clostridium thermocellum]	1.5	

	N	learest Neighbor		Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID	3.604.661	TY	1	<none></none>	<none></none>	<none></none>	
2593	M34661	Human chaperonin (HSP60) non- functional	1	NONE	NONL	STONE	
		pseudogene 3.					
2594	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	1723894	HYPOTHETICAL GTP-BINDING PROTEIN IN SEH1-PRP20 INTERGENIC REGION >gi 2131584 pir S 64106 hypothetical protein YGL099w - yeast (Saccharomyces cerevisiae) >gi 1322637 gnl PI D e243302 (Z72621) ORF YGL099w [Saccharomyces cerevisiae]	9e-015	
2595	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	0.16	
2596	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.4	
2597	U57715	Rattus norvegicus FGF receptor activating protein FRAG1 (FRAG1) mRNA, complete cds		1518609	(U57715) FGF receptor activating protein FRAG1 [Rattus norvegicus]	2e-088	
2598	Z64776	H.sapiens CpG DNA, clone 167d8, forward read cpg167d8.ft1b.	0.0002	1777782	(U52513) ISG family member [Homo sapiens]	2.4	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
2599	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA,	3e-008	<none></none>	<none></none>	<none></none>	
		complete cds					
2600	AF022158	Homo sapiens KRAB domain zinc finger protein	3e-010	2507553	ZINC FINGER PROTEIN 33A (ZINC FINGER PROTEIN KOX31) (KIAA0065) (HA0946) Kruppel-related. [Homo sapiens]	1e-016	
2601	Y07660	M.tuberculosis accBC gene	2e-068	465847	HYPOTHETICAL 66.5 KD PROTEIN F02A9.5 IN CHROMOSOME III >gi 280542 pir S2 8313 hypothetical protein F02A9.5 - Caenorhabditis elegans Genefinder; similar to Propionyl-CoA carboxylase beta chain; cDNA EST EMBL:M89018 comes from this gene; cDNA EST EMBL:D2806	8e-075	
2602	S51858	MO25 gene [mice, embryos, mRNA, 2322 nt]	0	547911	MO25 PROTEIN >gi 2143483 pir I5 7997 hypothetical calcium-binding protein - mouse protein [mice, embryos, Peptide, 341 aa] [Mus sp.]	e-119	
2603	AB018345	Homo sapiens mRNA for KIAA0802 protein, partial cds	e-131	3882325	(AB018345) KIAA0802 protein [Homo sapiens]	3e-053	

	<u> </u>	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2604	L41560	Homo sapiens (clones HGPCD2 and HGPCD15) pterin-4a- carbinolamine dehydratase (PCBD) gene, complete cds.	2e-005	<none></none>	<none></none>	<none></none>	
2605	AJ000041	Homo sapiens mRNA for HOXC11	e-180	987050	(X65335) lacZ gene product [unidentified cloning vector]	0.001	
2606	U55939	Expression vector pVP-Nco, complete sequence.	4e-043	987050	(X65335) lacZ gene product [unidentified cloning vector]	9e-009	
2607	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-014	124139	TRANS-ACTING TRANSCRIPTIO NAL PROTEIN ICP0 >gi 73901 pir WZ BE61 gene 61 protein - human herpesvirus 3 >gi 60050 (X04370) ORF 61 (AA1-467) [Human herpesvirus 3] >gi 228664 prf 18 08271A gene 61 protein	0.48	
2608	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
2609	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-013	3878130	(Z83112) predicted using Genefinder	9	
2610	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.3	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2611	D14965	C.elegans gene for alpha-2 tubulin, complete cds	3.7	<none></none>	<none></none>	<none></none>	
2612	Z61840	H.sapiens CpG DNA, clone 59g12, forward read cpg59g12.ft1a.	2e-080	3581872	(AL031541) putative integral membrane protein [Streptomyces coelicolor]	1.4	
2613	U59924	Sus scrofa nitric oxide synthase (NOS) mRNA, complete cds	1.1	<none></none>	<none></none>	<none></none>	
2614	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
2615	AF054625	Reporter vector pSRF-Luc, complete sequence	4e-065	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	
2616	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.1	
2617	AF031924	Homo sapiens homeobox transcription factor barx2	e-161	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	
2618	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
2619	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
2620		Reporter vector pCRE-Luc, complete sequence	1e-013	1065484	(U40415) similar to S. cerevisiae LAG1 (SP:P38703)	0.49	
2621	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete	1e-009	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		cds					
2622	AF013758	Homo sapiens polyadenylate binding protein- interacting protein-1 (PAIP1) mRNA, complete cds	0	3046900	(AF013758) polyadenylate binding protein- interacting protein-1 [Homo sapiens]	3e-072	
2623	D29808	Human mRNA for T-cell acute lymphoblastic leukemia associated antigen 1 (TALLA-1), complete cds	0.014	<none></none>	<none></none>	<none></none>	
2624	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	2690005	(AE000794) B. burgdorferi predicted coding region BBF30	7.6	
2625	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.041	<none></none>	<none></none>	<none></none>	
2626	Z12112	pWE15A cosmid vector DNA	2e-067	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-008	
2627	AB018326	Homo sapiens mRNA for KIAA0783 protein, complete cds	0	3882287	(AB018326) KIAA0783 protein [Homo sapiens]	1e-073	
2628	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.4	
2629	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-016	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.4	

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2630	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	1001632	(D64002) hypothetical protein	3.2	
2631	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	0.29	
2632	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
2633	X05167	Barley gene for thiol protease aleurain	0.13	1065515	(U40420) weak similarity to procollagen alpha chain 1(V) chain [Caenorhabditis elegans]	9e-018	
2634	Z96177	H.sapiens telomeric DNA sequence, clone 10QTEL040, read 10QTELOO040.s	2e-060	987050	(X65335) lacZ gene product [unidentified cloning vector]	5e-010	
2635	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	123432	ZERKNUELLT PROTEIN 1 (ZEN-1)	3.4	
2636	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	123432	ZERKNUELLT PROTEIN 1 (ZEN-1)	3.4	
2637	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.5	
2638	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.5	

		learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
2639	AF103734	Sindbis-like virus	3.5	<none></none>	<none></none>	<none></none>	
		YN87448,					
		complete genome		0.500.606	(4.0002525)	C- 010	
2640	M27280	H.influenzae lic-1	3.4	2529686	(AC002535)	6e-018	
		operon licA, licB,			putative G-beta- repeat containing		
		licC and licD			protein, 5' partial		
		genes, encoding outer membrane			[Arabidopsis		
		lipopolysaccharid			thaliana]		
		e phase variation,			,		
		complete cds.					
2641	AF103734	Sindbis-like virus	3.5	<none></none>	<none></none>	<none></none>	
20.11	711 103 / 5 .	YN87448,					
		complete genome					
2642	X05167	Barley gene for	0.13	1065515	(U40420) weak	9e-018	
		thiol protease			similarity to		
		aleurain			procollagen alpha		
					chain 1(V) chain		
					[Caenorhabditis		
				10.46000	elegans]	1 - 005	
2643	L76159	Homo sapiens	4e-032	1246233	(L76159) FRG1 gene product	1e-005	
		FRG1 mRNA,			[Homo sapiens]		
2644	AF086047	Complete cds. Homo sapiens	3e-008	628916	Delta-12	6	
2644	AFU8604/	full length insert	36-008	028910	desaturases -		
		cDNA clone			Anabaena		
		YX84A05			variabilis		
					desaturașe		
					[Anabaena		
					variabilis]		
2645	AF086136	Homo sapiens	4e-021	3849864	(AJ007629) pall	4.6	
		full length insert			protein		
		cDNA clone			[Emericella		
0.515	17004010	ZA89C06	1. 011	AIONES	nidulans] <none></none>	<none></none>	
2646	AB004818	Homo sapiens	1e-011	<none></none>	NONE	NOINE	
		mRNA for ENX- 2, complete cds					
2647	D87686	Homo sapiens	e-165	3540219	(D87686)	5e-054	
2047	D67080	mRNA for	0-103	3340217	KIAA0017 protein	30 05 1	
		KIAA0017			[Homo sapiens]		
		protein, complete			•		
1		cds					
2648	Z49218	S.cerevisiae	0.002	2984715	(AF053957)	0.33	
		chromosome XIII			dynamin		
		cosmid 7056			associated protein		
		<u></u>			isoform Dap160-1		

		learest Neighbor		Nearest Neighbor			
	`	astN vs. Genbank)			s. Non-Redundant Pro		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2649	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	868241	(U29488) C56C10.3 gene product [Caenorhabditis elegans]	7e-030	
2650	D38417	Mouse mRNA for arylhydrocarbon receptor, complete cds	3e-028	<none></none>	<none></none>	<none></none>	
2651	L29252	Human (clone D13-2) L-iditol-2 dehydrogenase gene, exon 4, exon 5, exon 6 and exon 7.	0.35	<none></none>	<none></none>	<none></none>	
2652	U29171	Human casein kinase I delta mRNA, complete cds >	3e-063	1176666	HYPOTHETICAL 139.1 KD PROTEIN C08B11.3 IN CHROMOSOME II >gi 3874171 gnl PI D e1343795 proteins; cDNA EST EMBL:T01154 comes from this gene; cDNA EST EMBL:T02016 comes from this gene; cDNA EST EMBL:D34307 comes from this gene; cDNA EST EMBL:D34307 comes from this	6.8	
2653	U63648	Mus musculus p160 myb- binding protein (P160) mRNA, complete cds	6e-058	2645205	(U63648) p160 myb-binding protein [Mus musculus]	2e-038	
2654	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.9	
2655	Y11740	H.sapiens whn gene, exon 1a and 1b	0.12	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2656	D26179	Rat mRNA for V-1 protein, complete cds		3879121	(Z70310) predicted using Genefinder; Similarity to Mouse ankyrin (PIR Acc. No. S37771); cDNA EST EMBL:T01923 comes from this gene; cDNA EST EMBL:D32335 comes from this gene; cDNA EST EMBL:D32723 comes from this gene; cDNA EST EMBL:D32723 comes from this gene; cDNA EST EMBL:D32723 comes from this gene; cDNA ES Genefinder; Similarity to M	8e-087	
2657	U67518	Methanococcus jannaschii section 60 of 150 of the complete genome	1.2	3876465	(Z81071) predicted using Genefinder; Similarity to Human small nuclear ribonucleoprotein E cDNA EST yk375g7.5 comes from this gene; cDNA EST yk435f5.3 comes from this gen	6e-011	
2658	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	<none></none>	<none></none>	<none></none>	
2659	U83176	Mus musculus ROSA 26 transcription AS ROSA26AS mRNA, complete cds	0	1778861	(U83176) ROSA26AS [Mus musculus]	e-101	
2660	AB018374	Mus musculus GARP34 mRNA, complete cds	2e-065	3724364	(AB018374) GARP34 [Mus musculus]	7e-010	
2661	AB018374	Mus musculus GARP34 mRNA, complete cds	2e-065	3724364	(AB018374) GARP34 [Mus musculus]	7e-010	

	N	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)			s. Non-Redundant Pr			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
2662	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.6		
2663	AL022168	Human DNA sequence from clone U247E12 on chromosome Xq22-23, complete sequence [Homo sapiens]	8e-008	<none></none>	<none></none>	<none></none>		
2664	M10277	Human cytoplasmic beta- actin gene, complete cds.	5e-063	<none></none>	<none></none>	<none></none>		
2665	D83769	Homo sapiens DNA, corresponding sequence for DHFR	5e-014	763429	(U22961) putative ORF; similar in part to the product encoded by human glycerol-3-phosphate dehydrogenase mRNA, GenBank Accession Number L34041; Method: conceptual translation supplied by author [Homo sapiens]			
2666	U15426	Human anonymous mRNA sequence with CCA repeat region.	3e-071	1065484	(U40415) similar to S. cerevisiae LAG1 (SP:P38703)	3e-015		
2667		Homo sapiens timing protein CLK-1 mRNA, complete cds	0	3811295	(AF032900) timing protein CLK-1; ubiquinone biosynthesis protein COQ7 [Homo sapiens]	3e-061		
2668	L39210	Homo sapiens inosine monophosphate dehydrogenase type II gene, complete cds	e-111	2887425	(AB007885) KIAA0425 [Homo sapiens]	3e-036		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2669	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7.00E-07	<none></none>	<none></none>	<none></none>	
2670	X93016	S.scrofa mRNA for cytosolic malic enzyme NADP-dependent	5e-045	101706	hypothetical protein 2 - yeast (Saccharomyces kluyveri) plasmid pSKL >gi 4870 (X54850) ORF 2, has similarity to DNA polymerase [Saccharomyces kluyveri]	7.7	
2671	J03068	Human DNF1552 (lung) mRNA, complete cds.	0.041	2414623	(Z99259) putative phosphotransferas e	7e-021	
2672	X81372	H.sapiens mRNA for biphenyl hydrolase-related protein	2e-016	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.0001	
2673	AB012130	Homo sapiens SBC2 mRNA for sodium bicarbonate cotransporter2, complete cds	0.00E+00	3097316	(AB012130) sodium bicarbonate cotransporter2 [Homo sapiens]	3e-045	
2674	D83769	Homo sapiens DNA, corresponding sequence for DHFR	5e-014	763429	(U22961) putative ORF; similar in part to the product encoded by human glycerol-3-phosphate dehydrogenase mRNA, GenBank Accession Number L34041; Method: conceptual translation supplied by author [Homo sapiens]		
2675	D38522	Human mRNA for KIAA0080 gene, partial cds	1e-022	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.002	
2676	D38522	Human mRNA for KIAA0080 gene, partial cds	1e-022	728831	!!!! ALU SUBFAMILY J WARNING	0.002	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins) ACCESSION DESCRIPTION P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION		PVALUE	
					ENTRY		
2677	AF072810	Homo sapiens transcription factor WSTF mRNA, complete cds	0	4049922	(AF072810) transcription factor WSTF [Homo sapiens]	1e-070	
2678	U41767	Human metargidin precursor mRNA, complete cds	e-130	1235674	(U41767) metargidin precursor [Homo sapiens]	1.00E-02	
2679	L81613	Homo sapiens (subclone 4_c7 from P1 H17) DNA sequence	0.38	<none></none>	<none></none>	<none></none>	
2680	M68841	Human L1 repetitive sequence with a region homologous to a mouse ORF.	9.00E-30	106322	hypothetical protein (L1H 3' region) - human	8e-008	
2681	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0005	<none></none>	<none></none>	<none></none>	
2682	D87973	Mus musculus Impact mRNA, complete cds	0	4038076	(D87973) Impact [Mus musculus]	1e-095	
2683	M69175	Human H-protein mRNA, complete cds.	2e-017	<none></none>	<none></none>	<none></none>	
2684	Z80361	H.sapiens HLA- DRB pseudogene, repeat region;	1e-082	1706108	MITOCHONDRI AL CARNITINE O- PALMITOYLTR ANSFERASE I, LIVER ISOFORM (CPT I) (CPTI-L) carnitine palmitoyltransfera se I [Homo sapiens] I [Homo sapiens]	0.67	

SEQ ID		Nearest Neighbor astN vs. Genbank)			Nearest Neighbor		
	ACCESSION			(BlastX vs. Non-Redundant Proteins)			
		DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2685	AF017044	Dictyostelium discoideum LTR- retrotransposon Skipper, partial genomic sequence, 3' end	0.014	<none></none>	<none></none>	<none></none>	
2686	U40825	Mus musculus WW-domain binding protein 1 mRNA, complete cds	e-118	1777577	(U40825) WW- domain binding protein 1 [Mus musculus]	2.00E-29	
2687	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	2281149	(U58553) maturase [Carum carvi]	4.6	
2688	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-008	3328840	(AE001314) Putative outer membrane protein A [Chlamydia trachomatis]	5.8	
2689	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
2690	AB012130	Homo sapiens SBC2 mRNA for sodium bicarbonate cotransporter2, complete cds	0.00E+00	3097316	(AB012130) sodium bicarbonate cotransporter2 [Homo sapiens]	3e-045	
2691	X69516	H.sapiens gene for folate receptor	3e-008	<none></none>	<none></none>	<none></none>	
2692	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5.00E-04	1203965	(L42379) bonederived growth factor [Homo sapiens]	0.17	
2693	Z15027	H.sapiens HLA class III DNA	3.00E-07	728836	!!!! ALU SUBFAMILY SP WARNING ENTRY	3.6	
2694	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	2e-006	<none></none>	<none></none>	<none></none>	
2695	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	2e-006	<none></none>	<none></none>	<none></none>	

	<u> </u>	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2696	X77775	G.gallus Gal beta 1, 3 GalNAc- specific GalNAc alpha 2, 6- sialyltransferase mRNA.	1e-022	3873839	(Z81029) W05H12.2 [Caenorhabditis elegans] >gi 3880545 gnl PI D e1350077 (Z82072) W05H12.2	5.9	
2697	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	2281149,	(U58553) maturase [Carum carvi]	4.6	
2698	U33005	Mus musculus tbc1 mRNA, complete cds. > :: gb 186429 186429 Sequence 1 from patent US 5700927	3e-093	3893077	(Y17923) lyncein [Bos taurus]	1e-040	
2699	U74651	Human DNA polymerase gamma (polg) gene, promoter region and partial cds	1e-022	113667	!!!! ALU CLASS B WARNING ENTRY !!!!	0.002	
2700	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-009	3064257	(AF043899) amphiphysin IIc1 [Homo sapiens]	0.87	
2701	U43893	Mus musculus ATP synthase gamma-subunit gene, nuclear gene encoding a mitochondrial protein, partial cds	0.005	3929529	(AF034611) intrinsic factor- B12 receptor precursor; cubilin [Homo sapiens]	0.67	
2702	U43893	Mus musculus ATP synthase gamma-subunit gene, nuclear gene encoding a mitochondrial protein, partial cds	0.005	3929529	(AF034611) intrinsic factor- B12 receptor precursor; cubilin [Homo sapiens]	0.67	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID				110554	AMPHIREGULIN	4e-041	
2703	M30704	Human amphiregulin (AR) mRNA, complete cds, clones lambda- AR1 and lambda- AR2.	0	113754	PRECURSOR (AR)		
2704	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-010	<none></none>	<none></none>	<none></none>	
2705	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.2	
2706	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	2832664	(AL021710) pollen-specific protein - like [Arabidopsis thaliana]	8e-020	
2707	U00684	Human unknown mRNA.	2e-038	2500412	30S RIBOSOMAL PROTEIN S6 Mycoplasma pneumoniae (SGC3) (ATCC 29342) >gi 1674305 similar to Swiss- Prot Accession Number P02358, from E. coli [Mycoplasma pneumoniae]	1.3	
2708	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5e-015	108693	glutamic acid-rich protein, retinal - bovine taurus]	0.067	
2709	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	79703	hypothetical 32K protein (frxC 5' region) - Synechocystis sp. (PCC 6803) >gi 217091 gnl PI D d1001745	0.8	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2710	AF083395	Homo sapiens phospholipase A2-activating protein mRNA, complete cds	e-175	4106818	(AF083395) phospholipase A2- activating protein [Homo sapiens]	4e-039	
2711	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-011	<none></none>	<none></none>	<none></none>	
2712	AB019488	Homo sapiens DNA for TRKA, exon 17 and complete cds	0	37403	(X03541) trk gene product (aa 1-641) [Homo sapiens]	1e-032	
2713	X62570	H.sapiens mRNA for IFP53	e-105	32709	(X62570) IFP53 [Homo sapiens]	6e-033	
2714	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.005	1170056	GENERAL SECRETION PATHWAY PROTEIN F	4.3	
2715	AF031924	Homo sapiens homeobox transcription factor barx2	e-161	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	
2716	AF031924	Homo sapiens homeobox transcription factor barx2	e-161	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	
2717	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.4	
2718	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.4	
2719	L20826	Human I-plastin mRNA, complete cds.	e-163	2493466	I-PLASTIN (INTESTINE- SPECIFIC PLASTIN) >gi 1362892 pir A 56536 plastin, intestine-specific- human >gi 405230 (L20826) I-plastin	6e-069	

	, D	Nearest Neighbor		Nearest Neighbor			
	(Bl	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2720	Z54386	H.sapiens CpG DNA, clone 10g3, forward read cpg10g3.ft1a	7e-059	1788180	(AE000281) biotin sulfoxide reductase 2 [Escherichia coli]	5.8	
2721	AF086201	Homo sapiens full length insert cDNA clone ZC42G09	1e-085	2564332	(AB006630) KIAA0292 [Homo sapiens]	5.4	
2722	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	0.12	
2723	AJ006267	Homo sapiens mRNA for ClpX- like protein	0	3688380	(AJ006267) ClpX- like protein [Homo sapiens]	1e-091	
2724	AF064801	Homo sapiens multiple membrane spanning receptor TRC8 (TRC8) mRNA, complete cds	0	3395787	(AF064801) multiple membrane spanning receptor TRC8	e-123	
2725	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	2599526	(AF029331) immunoglobulin heavy chain V region [Homo sapiens]	4.2	
2726	Y08013	S.salar DNA segment containing GT repeat	0.006	<none></none>	<none></none>	<none></none>	
2727	Y08013	S.salar DNA segment containing GT repeat	0.006	<none></none>	<none></none>	<none></none>	
2728	AE000971	Archaeoglobus fulgidus section 136 of 172 of the complete genome	0.041	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	(B)	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2729	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-007	1170586	RAS GTPASE-ACTIVATING-LIKE PROTEIN IQGAP1 (P195) (KIAA0051) >gi 627594 pir A5 4854 Ras GTPase activating-related protein - human sapiens] >gi 536844 (L33075) ras GTPase-activating-like protein [Homo sapiens]	9e-011	
2730	M60858	Human nucleolin gene, complete cds.	e-129	<none></none>	<none></none>	<none></none>	
2731	M85145	Human tumor necrosis factor receptor, 3' flank.	2e-007	<none></none>	<none></none>	<none></none>	
2732	M85145	Human tumor necrosis factor receptor, 3' flank.	2e-007	<none></none>	<none></none>	<none></none>	
2733	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-013	<none></none>	<none></none>	<none></none>	
2734	L07063	Mus musculus FKBP65 binding protein mRNA, complete cds	6e-089	2137294	FKBP65 binding protein - mouse >gi 894162	6e-024	
2735	X63432	H.sapiens ACTB mRNA for mutant beta-actin	e-112	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-014	
2736	AF083395	Homo sapiens phospholipase A2-activating protein mRNA, complete cds	0	4106818	(AF083395) phospholipase A2- activating protein [Homo sapiens]	1e-094	
2737	AJ012449	Homo sapiens mRNA for NS1- binding protein	3e-009	3165570	(AF067946) similar to Drosophila ring canal protein	4e-032	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2738	M27878	Human DNA binding protein (HPF2) mRNA, complete cds.	3e-063	3702137	(AL031393) dJ733D15.1 (Zinc- finger protein) [Homo sapiens]	1e-040	
2739	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
2740	Y15230	Homo sapiens pygl gene, exon 5 and partial intron 4 and 5	e-166	3170407	(AF046798) glycogen phosphorylase [Homo sapiens]	1e-044	
2741	Z96177	H.sapiens telomeric DNA sequence, clone 10QTEL040, read 10QTELOO040.s eq		987050	(X65335) lacZ gene product [unidentified cloning vector]	2e-005	
2742	M90058	Human serglycin gene, exons 1,2, and 3.	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	8.8	
2743	X69878	H.sapiens Flt4 mRNA for transmembrane tyrosine kinase	2e-088	<none></none>	<none></none>	<none></none>	
2744	X69878	H.sapiens Flt4 mRNA for transmembrane tyrosine kinase	2e-088	<none></none>	<none></none>	<none></none>	
2745	AB007923	Homo sapiens mRNA for KIAA0454 protein, partial cds	0	3413870	(AB007923) KIAA0454 protein [Homo sapiens]	1e-098	
2746	AF042181	Homo sapiens testis-specific Y- encoded-like protein (TSPYL) mRNA, partial cds	2e-047	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3	
2747	AL021173	Caenorhabditis elegans cosmid VK10D6R, complete sequence [Caenorhabditis elegans]	1.2	<none></none>	<none></none>	<none></none>	
2748	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	0.12	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2749	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-008	<none></none>	<none></none>	<none></none>	
2750	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-008	<none></none>	<none></none>	<none></none>	
2751	M22970	Human pancreatic phospholipase A- 2 (PLA-2) gene, exons 1 to 3.	1e-032	113671	!!!! ALU CLASS F WARNING ENTRY !!!!	3e-006	
2752	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.2	
2753	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-011	3219914	HYPOTHETICAL 16.8 KD PROTEIN C30D10.04 IN CHROMOSOME II >gi 2276353 gnl PI D e330328 pombe]		
2754	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	3875246	(Z81490) similar to WD domain, Gbeta repeats (2 domains); cDNA EST EMBL:T00482 comes from this gene; cDNA EST EMBL:T00923 comes from this gene; cDNA EST yk449d4.3 comes from this gene; cDNA EST yk449d4.5 comes from this gene;	6e-078	

	7	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)	,	(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
2755	D79205	Human mRNA for ribosomal protein L39, complete cds	1e-086	1173044	60S RIBOSOMAL PROTEIN L39 norvegicus] >gi 1373419 (U57846) ribosomal protein L39 ribosomal protein L39 [Homo sapiens]	4e-009		
2756	AB014591	Homo sapiens mRNA for KIAA0691 protein, complete cds	0	3327196	(AB014591) KIAA0691 protein [Homo sapiens]	1e-047		
2757	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-012	115409	CUTICLE COLLAGEN ROL-6 elegans] >gi 3879235 gnl PI D e1348932 (Z66499) similar to cuticle collagen ROL-6; cDNA EST cm10c4 comes from this gene; cDNA EST EMBL:M88874 comes from this gene; cDNA EST yk265e2.3 comes from this gene; cDNA EST yk265e2.5 comes fro	0.031		
2758	U78096	Human macrophage colony stimulating factor receptor (c-fms) gene, exon 1A, 2 and partial cds	4e-012	126296	LINE-1 REVERSE TRANSCRIPTAS E HOMOLOG protein [Nycticebus coucang]	0.0005		
2759	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	<none></none>	<none></none>	<none></none>		

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2760	M27878	Human DNA binding protein (HPF2) mRNA, complete cds.	3e-063	3702137	(AL031393) dJ733D15.1 (Zinc- finger protein) [Homo sapiens]	1e-040	
2761	U43076	Mus musculus cdc37 homolog mRNA, complete cds	2e-017	755484	(U20281) cell division cycle control protein 37 [Gallus gallus]	8e-022	
2762	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.2	
2763	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	1171883	SODIUM- INDEPENDENT ORGANIC ANION TRANSPORTER (ORGANIC ANION TRANSPORTING POLYPEPTIDE) anion - rat >gi 410311 (L19031) oatp [Rattus norvegicus]	2e-036	
2764	X54452	D.discoideum culmination spiA (Dd31) gene	3.3	<none></none>	<none></none>	<none></none>	
2765	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	<none></none>	<none></none>	<none></none>	
2766	AF053698	Reporter vector pAP1-Luc, complete sequence	3e-019	987050	(X65335) lacZ gene product [unidentified cloning vector]	0.2	
2767		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-012	3582428	(AB017257) glycocyamine kinase beta chain [Neanthes diversicolor]	4.3	
2768		<none></none>	<none></none>		<none></none>	<none></none>	
2769	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial	6e-006	3511122	(AF060503) zinc finger protein [Homo sapiens]	5.3	

	7	Nearest Neighbor		Nearest Neighbor			
	(B)	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		cds					
2770	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	<none></none>	<none></none>	<none></none>	
2771	Ù95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	<none></none>	<none></none>	<none></none>	
2772	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
2773	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-013	804788	(M13002) 2855 is the position of the first start codon in ORF 2; putative [Mus musculus]	0.64	
2774	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	8e-008	<none></none>	<none></none>	<none></none>	
2775	M86526	Rat proline-rich protein (PRP) gene, 5' end, and containing several Alu-like repetitive elements.	0.37	<none></none>	<none></none>	<none></none>	
2776	Z22923	M.musculus alpha2 (IX) collagen gene, complete CDS.	0.002	<none></none>	<none></none>	<none></none>	
2777	Z22923	M.musculus alpha2 (IX) collagen gene, complete CDS.	0.002	<none></none>	<none></none>	<none></none>	
2778	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.4	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
2779	Z74035	Caenorhabditis elegans cosmid F47G9, complete sequence [Caenorhabditis elegans]	3.4	2879805	(AL021813) hypothetical protein	5.7	
2780	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	<none></none>	<none></none>	<none></none>	
2781	AG001356	Homo sapiens genomic DNA, 21q region, clone: 9H11BG25	2e-015	<none></none>	<none></none>	<none></none>	
2782	D83006	Saccharomyces cerevisiae MNN4 gene, complete cds	1.2	<none></none>	<none></none>	<none></none>	
2783	Z59640	H.sapiens CpG DNA, clone 167g11, forward read cpg167g11.ft1b.	0.12	<none></none>	<none></none>	<none></none>	
2784	AF049069	Pinus radiata PRE87 mRNA, complete cds	1.1	1518141	(U66568) myocyte enhancer factor 2A MEF2A [Danio rerio]	3.1	
2785	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.4	
2786	AF031931	Hydra oligactis cyclic GMP- dependent protein kinase (hyGK) mRNA, complete cds	0.13	<none></none>	<none></none>	<none></none>	
2787	Z96177	H.sapiens telomeric DNA sequence, clone 10QTEL040, read 10QTELOO040.s eq	3e-041	987050	(X65335) lacZ gene product [unidentified cloning vector]	0.015	

		Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)			s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2788	L48716	Homo sapiens galactose-1- phosphate uridyl transferase (GALT) mutant F117S gene, exons 3 and 4	1.1	77657	hypothetical 30.1K protein - Pseudomonas aeruginosa	0.095	
2789	U73902	Mus musculus emerin (Sta) mRNA, complete cds	0.37	529773	(U06752) Heterodimeric complex composed of a mucin subunit, ASGP-1, which is predominantly Oglycosylated, and a cysteine-rich transmembrane subunit, ASGP-2, which is predominantly Nglycosylated [Rattus norvegicus]	0.009	
2790	X54171	H.sapiens NG2-6 DNA	4e-021	<none></none>	<none></none>	<none></none>	
2791	M30519	Mouse mammary tumor virus gag gene, 3' end, pol gene, 5' end.	0.12	1262926	(U51903) RasGAP-related protein [Homo sapiens]	4.3	
2792	AJ223355	Rattus norvegicus mRNA for mitochondrial dicarboxylate carrier	0.38	128059	NEGATIVE FACTOR (F- PROTEIN) (27 KD PROTEIN) (3'ORF) >gi 77283 pir S07 993 nef protein - simian immunodeficiency virus SIVsm (isolate F236) immunodeficiency virus]	2	
2793	AF086022	Homo sapiens full length insert cDNA clone YW23E02	6e-005	3402679	(AC004697) unknown protein [Arabidopsis thaliana]	9e-016	
2794	U47322	Cloning vector DNA, complete sequence.	9e-010	<none></none>	<none></none>	<none></none>	

1.		Nearest Neighbor		Nearest Neighbor			
0.70		lastN vs. Genbank)	T 50 T 5 T 7 T 7 T 7 T 7 T 7 T 7 T 7 T 7 T 7	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2795	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	3873667	(Z71178) similar to collagen	0.093	
2796	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	2745961	(U51869) Bcd orf2 [Homo sapiens]	0.47	
2797	AF041209	Homo sapiens midline 1 fetal kidney isoform 2	0.0002	<none></none>	<none></none>	<none></none>	
2798	AF092564	Homo sapiens chromosome- associated protein-C	5e-056	4092846	(AB019987) chromosome- associated polypeptide-C [Homo sapiens]	3e-017	
2799	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	<none></none>	<none></none>	<none></none>	
2800	M95623	Homo sapiens hydroxymethylbil ane synthase gene, complete cds.	0.005	4007760	(AL034433) importin alpha subunit	4.2	
2801	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
2802	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	1065945	(U40799) coded for by C. elegans cDNA yk28f2.3; coded for by C. elegans cDNA yk12c10.3; coded for by C. elegans cDNA yk5a12.3; coded for by C. elegans cDNA yk49a8.3; coded for by C. elegans cDNA yk12c10.5; coded for by C. elegans cDNA yk12c10.5; coded for by C. elegans cDNA yk28f2	0.12	

	Ň	learest Neighbor		Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2803	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete	0.04	<none></none>	<none></none>	<none></none>	
2804	M74558	Human SIL mRNA, complete cds. > :: gb G28581 G285 81 human STS SHGC-35335.	e-126	<none></none>	<none></none>	<none></none>	
2805	M72885	Human GOS2 gene, 5' flank and cds.	0.36	3873821	(Z68213) cDNA EST yk266c4.5 comes from this gene; cDNA EST yk266c4.3 comes from this gene	1.8	
2806	U27341	Bos taurus endothelin converting enzyme-2 Sequence 1 from patent US 5736376	6e-078	2136744	endothelin converting enzyme-2 - bovine	3e-028	
2807	U36756	Mus musculus thrombin receptor (Cf2r) gene, exon	0.013	<none></none>	<none></none>	<none></none>	
2808	AJ003209	Human immunodeficienc y virus type 1 mRNA for reverse transcriptase, isolate H-20, partial	0.12	<none></none>	<none></none>	<none></none>	
2809	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	<none></none>	<none></none>	<none></none>	
2810	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-009	1272701	(L11900) cytochrome b [Cratogeomys bulleri]	9.3	

4	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID			2 222	ALONES	AIONES	<none></none>	
2811	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	<none></none>	<none></none>		
2812	AB006572	Homo sapiens RMP mRNA for RPB5 meidating protein, complete cds	0	3970833	(AB006572) RPB5 meidating protein [Homo sapiens]	5e-037	
2813	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	1109865	(U41540) coded for by C. elegans cDNA yk42d12.5; coded for by C. elegans cDNA yk27e10.5; coded for by C. elegans cDNA cm08h6; coded for by C. elegans cDNA yk88e12.5; coded for by C. elegans cDNA yk42d12.3; coded for by C. elegans cDNA yk27e1	2e-009	
2814	Z26259	H.sapiens isoform 1 gene for L-type calcium channel, exon 4		3426264	(AF037269) cell division protein [Mycobacterium smegmatis]	0.47	
2815		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	2358285	(AF010403) ALR [Homo sapiens]	0.27	
2816		Homo sapiens chromosome 5, P1 clone 1209C1 (LBNL H104), complete sequence [Homo sapiens]	2e-006	<none></none>	<none></none>	<none></none>	
2817	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.1	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2818	U43626	Human chromosome 15q11-q13 putative DNA replication origin in the g- aminobutyric acid receptor b3 and a5 gene cluster	2e-018	2197085	(AF003535) ORF2-like protein [Homo sapiens]	0.0002	
2819	Z96402	H.sapiens telomeric DNA sequence, clone 18QTEL022, read 18QTELOO022.s eq		386792	intercellular adhesion molecule 2 (ICAM-2) [Homo sapiens]	9.2	
2820	U43626	Human chromosome 15q11-q13 putative DNA replication origin in the g- aminobutyric acid receptor b3 and a5 gene cluster	2e-018	2197085	(AF003535) ORF2-like protein [Homo sapiens]	0.0002	
2821	U66534	Human beta4- integrin (ITGB4) gene, exon 14,15,16,17 and 18	0.12	<none></none>	<none></none>	<none></none>	
2822	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.2	
2823	AC001462	Homo sapiens (subclone 2_h10 from BAC H107) DNA sequence		2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	7.1	
2824	AE000464	Escherichia coli K-12 MG1655 section 354 of 400 of the complete genome		3879850	(Z81592) predicted using Genefinder	2e-039	
2825	AB018304	Homo sapiens mRNA for KIAA0761 protein, partial cds	3e-009	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2826	AL008982	Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS *** from contig 3-52, complete sequence	3.2	3880930	(AL021481) similar to Phosphoglucomuta se and phosphomannomut ase phosphoserine; cDNA EST EMBL:D36168 comes from this gene; cDNA EST EMBL:D70697 comes from this gene; cDNA EST yk373h9.5 comes from this gene; cDNA EST yk373h9.5 comes		
2827	Z 54196	S.cereale DNA for repeat unit (D1100 family)	0.36	2500714	HYPOTHETICAL 35.0 KD PROTEIN F48E8.1 IN CHROMOSOME III >gi 746485 (U23514) similar to antigen domain of venom allergen (SP:VA52_DOLM A, P10736) and to antigen 5 (PIR:A37329) [Caenorhabditis elegans]	4.1	
2828	Z95979	Homo sapiens hRED1 gene, exons 7, 8, 9 and 10	7e-017	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	0.002	
2829	Z15030	H.sapiens gene for ventricular myosin light chain 2 > :: gb L01652 HUM VMLC Human ventricular myosin light chain 2 gene, seven exons.	5e-024	565265	(M76741) biliary glycoprotein [Homo sapiens]	9.2	
2830	U56440	Human His-1 gene sequence	8e-007	<none></none>	<none></none>	<none></none>	

4		Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
2831	AF009941	Tomocichla tuba	1.2	<none></none>	<none></none>	<none></none>	
		cytochrome b (cytb) gene,					
		mitochondrial					
		gene encoding					
		mitochondrial					
		protein, complete					
		cds			CONTRACTOR OF THE PROPERTY OF	1	
2832	X68011	H.sapiens ZNF81	3e-030	1731442	ZINC FINGER PROTEIN 81	1e-020	
		gene			human (fragment)	!	
					>gi 454325	1	
					(X68011) ZNF81		
					gene product		
2833	U36499	Human	1e-020	<none></none>	<none></none>	<none></none>	
		lymphoid-specific SP100 homolog					
		(LYSP100-A)					
		mRNA, complete					
		cds					
2834	Z60692	H.sapiens CpG	3e-059	<none></none>	<none></none>	<none></none>	
		DNA, clone 31f7,					
		reverse read cpg31f7.rt1a.					
2835	X92485	P.vivax pval	0.0002	<none></none>	<none></none>	<none></none>	
	1132.00	gene	0.0002	1,01,2	110112	1,01,12	
2836	U95102	Xenopus laevis	0.0005	576257	Prostatic Acid	3e-009	
		mitotic			Phosphatase .		
		phosphoprotein 90 mRNA,			(E.C.3.1.3.2) Complexed With		
		complete cds			Tartaric Acid		
		compiete eas			>gi 576258 pdb 1R		
					PT Prostatic Acid		
					Phosphatase		
					(E.C.3.1.3.2) Complexed With		
					Complexed with Vanadate		
					- minner	<u> </u>	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2837	U72372	Scandia geniculata 18S ribosomal RNA and 25S ribosomal RNA genes, partial sequence, and internal transcribed spacer 1, 5.8S ribosomal RNA gene and internal transcribed spacer 2, complete sequence		<none></none>	<none></none>	<none></none>	
2838	D49425	Anabaena variabilis rbpD gene for RNA- binding protein, complete cds	3.2	<none></none>	<none></none>	<none></none>	
2839	X95844	S.cerevisiae POP3 gene	3.5	<none></none>	<none></none>	<none></none>	
2840	AE001425	Plasmodium falciparum chromosome 2, section 62 of 73 of the complete sequence	0.041	3880909	(AL032636) Y40B1B.3 [Caenorhabditis elegans]	5.5	
2841	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>	
2842	X69064	M.musculus Ank- 1 mRNA for erythroid ankyrin		<none></none>	<none></none>	<none></none>	
2843	U61950	Caenorhabditis elegans cosmid C45E5	0.13	<none></none>	<none></none>	<none></none>	
2844	U73332	Human non- coding genomic sequence upstream from unique L0 sequence in the alpha-globin gene cluster	1e-010	<none></none>	<none></none>	<none></none>	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID					ALONES	AIONES	
2845	U21051	Human G protein-coupled receptor (GPR4) gene, complete cds.	0.13	<none></none>	<none></none>	<none></none>	
2846	X57921	O.sativa random single-copy DNA fragment 12RG214R	4.1	<none></none>	<none></none>	<none></none>	
2847	AF037219	Homo sapiens PIX1 mRNA sequence	0.043	<none></none>	<none></none>	<none></none>	
2848	M55124	Human cystic fibrosis transmembrane conductance regulator (CFTR) gene, exon 17b	0.005	<none></none>	<none></none>	<none></none>	
2849	AF035527	Mus musculus EHF (Ehf) mRNA, complete cds	e-164	3138930	[Mus musculus]	5e-084	
2850	AF052695	Rattus norvegicus cell cycle protein p55CDC gene, complete cds	3.7	2894379	(Y14573) ring finger protein [Hordeum vulgare]	8.2	
2851	<none></none>	<none></none>	<none></none>	3327112	(AB014549) KIAA0649 protein [Homo sapiens]	3.8	
2852	M34664	Human chaperonin (HSP60) mRNA, complete cds.	0	2501737	TRANSCRIPTIO NAL ACTIVATOR PROTEIN ACU- 15 >gi 1922895 gnl PI D e308394 (Y11565) transcriptional activator protein [Neurospora crassa]		
2853	D49701	Aspergillus oryzae niaD gene for nitrate reductase, complete cds	0.042	3879556	(Z70756) T06E4.11 [Caenorhabditis elegans]	0.5	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2854	AF016266	Homo sapiens TRAIL receptor 2 mRNA, complete cds	1e-010	134846	SMALL PROLINE-RICH PROTEIN II rich protein [Homo sapiens]	1.5	
2855	U44862	Human Down Syndrome region of chromosome 21, clone A11E6- 2B6.	1.2	<none></none>	<none></none>	<none></none>	
2856	X14503	Chlamydomonas eugametos petD gene for cytochrome b6/f complex subunit IV	0.13	<none></none>	<none></none>	<none></none>	
2857	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	3228515	(U70256) SomA [Synechococcus PCC6301]	4.6	
2858	M25534	Chicken actin- capping protein (CapZ 36/32) alpha subunit mRNA, complete cds.	0.41	<none></none>	<none></none>	<none></none>	
2859	X84372	D.melanogaster lethal(3)73Ah gene	1.1	<none></none>	<none></none>	<none></none>	
2860	AF053551	Homo sapiens metaxin 2 (MTX2) mRNA, nuclear gene encoding mitochondrial protein, complete cds	0	3283049	(AF053551) metaxin 2 [Homo sapiens]	2e-089	
2861	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-005	3877358	(Z66520) similar to RBB3 like protein; cDNA EST EMBL:C08891 comes from this gene; cDNA EST EMBL:C09371 comes from this gene; cDNA EST yk468f10.5 comes	3e-005	

		learest Neighbor		Nearest Neighbor			
	(BlastN vs. Genbank)			(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					from this gene [Caenorhabditis elegans]		
2862	AB002450	Homo sapiens mRNA from chromosome 5q21-22, clone:A3-A	2e-014	3790760	(AF099922) No definition line found [Caenorhabditis elegans]	2.5	
2863	AF053698	Reporter vector pAP1-Luc, complete sequence	1e-009	<none></none>	<none></none>	<none></none>	
2864	AF045086	Drosophila prosaltans 14045- 0901.4 cytochrome oxidase II (COII) gene, mitochondrial gene encoding mitochondrial protein, complete cds	0.005	<none></none>	<none></none>	<none></none>	
2865	Y09312	C.botulinum HA-70 gene (partial) and HA-17 gene	0.002	1171601	(X95276) rps8 [Plasmodium falciparum]	5.7	
2866	AJ001597	Homo sapiens gene encoding cAMP-dependent protein kinase gamma isoform	0.005	1869883	(Z86099) RS1 [human herpesvirus 2] herpesvirus 2]	0.52	
2867	AF022962	Mus musculus Sec8 mRNA, complete cds	1.1	<none></none>	<none></none>	<none></none>	
2868	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0005	2499622	PROBABLE SERINE/THREO NINE-PROTEIN KINASE YOL113W (PROTEIN KINASE 75490	3.5	

		Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					D)		
2869	AJ005262	Dictyostelium discoideum gene encoding a novel glycoprotein	0.12	<none></none>	<none></none>	<none></none>	
2870	U08214	Rattus sp. DNA binding protein (URE-B1) mRNA, complete cds.	0.12	4033834	(AJ009556) cytoskeleton assembly control protein Sla2p [Candida albicans]	0.13	
2871	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
2872	M31061	Human ornithine decarboxylase gene, complete cds.	2e-015	3808095	(Y08560) SCO- spondin [Bos taurus]	0.098	
2873	U21914	Human duplicate spinal muscular atrophy mRNA, clone 5G7, partial cds.	0.002	<none></none>	<none></none>	<none></none>	
2874	<none></none>	<none></none>	<none></none>	1228047	(D83782) the KIAA0199 gene is expressed ubiquitously.; the KIAA0199 protein shows similarity to sea urchin hydroxymethylglut alyl-CoA reductase, and retains 8 hydrophobic domains. [Homo sapiens]	2.5	
2875	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	4105505	(AF046914) multiple inositol polyphosphate phosphatase	5.6	
2876	Z96210	H.sapiens telomeric DNA sequence, clone 12PTEL057, read 12PTELOO057.s eq	0.014	2347056	(AJ000085) Nedd4 protein [Xenopus laevis]	5.8	

	Ŋ	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2877	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	2133693	masquerade precursor - fruit fly (Drosophila melanogaster) >gi 665545 (U18130) masquerade [Drosophila melanogaster] >gi 1095942 prf 2 110286A masquerade gene	1.2	
2878	X54252	C. elegans complete mitochondrial genome	0.38	<none></none>	<none></none>	<none></none>	
2879	S81913	adrenocorticotrop in receptor [Papio anubis=baboons, adrenal, mRNA Partial, 426 nt]	1.2	<none></none>	<none></none>	<none></none>	
2880	X65997	M.musculus c-kit mRNA for truncated tyrosine-kinase	0.13	<none></none>	<none></none>	<none></none>	
2881	AE000588	Helicobacter pylori section 66 of 134 of the complete genome	1.1	<none></none>	<none></none>	<none></none>	
2882	U64861	Caenorhabditis elegans cosmid C47D2.	0.12	<none></none>	<none></none>	<none></none>	
2883	U23173	Caenorhabditis elegans cosmid K07E1	0.37	2854192	(AF045645) contains similarity to microsomal triglyceride transfer proteins [Caenorhabditis elegans]	7.2	
2884	AB014579	Homo sapiens mRNA for KIAA0679 protein, partial cds	0	3327172	(AB014579) KIAA0679 protein [Homo sapiens]	2e-053	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID				450505	(7700445)	0.17	
2885	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	1707032	(U80445) coded for by C. elegans cDNA yk13g5.3; coded for by C. elegans cDNA yk21g6.3; coded for by C. elegans cDNA CEMSE18F; coded for by C. elegans cDNA yk126b1.3; coded for by C. elegans cDNA yk65h8.3; coded for by C.	0.17	
2006	722705	H.sapiens	6e-005	<none></none>	elegans cDNA yk65h8 <none></none>	<none></none>	
2886	Z22795	microsatellite repeat.					
2887	AE001061	Archaeoglobus fulgidus section 46 of 172 of the complete genome	1.1	3738162	(AL031856) putative involvement in protein glycosylation in the golgi [Schizosaccharom yces pombe]	2.4	
2888	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	<none></none>	<none></none>	<none></none>	
2889	Z96643	H.sapiens telomeric DNA sequence, clone 5QTEL064, read 5QTELOO064.se	0.0005	1363732	probable membrane protein YLR454w - yeast	4	
2890	Z96643	H.sapiens telomeric DNA sequence, clone 5QTEL064, read 5QTELOO064.se q	0.0005	1363732	probable membrane protein YLR454w - yeast	4	

	<u>N</u>	learest Neighbor		Nearest Neighbor			
-	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2891	X80169	M.musculus mRNA for 200 kD protein	e-177	1717793	PROTEIN TSG24 (MEIOTIC CHECK POINT REGULATOR) >gi 1083553 pir A 55117 tsg24 protein - mouse	5e-069	
2892	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-009	3832555	(AF077439) immunoglobulin heavy chain variable region	4.4	
2893	AC002359	Homo sapiens Xp22 Cosmid U239B3 (from Lawrence Livermore X library) complete sequence [Homo sapiens]	2e-007	3599342	(AF081112) ORF2 [Mus musculus domesticus]	0.61	
2894	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-011	3123058	HYPOTHETICAL WD-REPEAT PROTEIN SLL0163 > gi 1001440 gnl PI D d1010715 (D63999) beta transducin-like protein [Synechocystis sp.]	0.001	
2895	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.2	
2896	Z46940	H.sapiens PRM1 gene, PRM2 gene and TNP2 gene	0.013	<none></none>	<none></none>	<none></none>	
2897	Z47735	H.sapiens NFKB1 gene, exons 11 & 12	2e-008	<none></none>	<none></none>	<none></none>	
2898	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.004	2224611	(AB002333) KIAA0335 [Homo sapiens]	4	

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2899	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
2900	X00367	Chlamydomonas chloroplast DNA region with ARS element 03 (ARS = autonomously replicating sequence)	0.12	<none></none>	<none></none>	<none></none>	
2901	U41222	Dictyostelium discoideum RacE (racE) gene, complete cds	0.35	<none></none>	<none></none>	<none></none>	
2902	AB007504	Triticum aestivum TaMADS#11 mRNA for MADS box transcription factor, complete cds	0.042	<none></none>	<none></none>	<none></none>	
2903	X65319	Cloning vector pCAT-Enhancer	7e-069	987050	(X65335) lacZ gene product [unidentified cloning vector]	7e-011	
2904	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0005	3924670	(AC004990) supported by Genscan and several ESTs: C83049	6e-042	
2905	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.041	2132051	hypothetical protein YOR083w - yeast	3.3	
2906		pWE15A cosmid vector DNA	6e-068	987050	(X65335) lacZ gene product [unidentified cloning vector]	2e-009	
2907	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	2995374	(AL022245) hypothetical protein	5e-005	

		Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2908	U47322	Cloning vector DNA, complete sequence.	3e-009	<none></none>	<none></none>	<none></none>	
2909	X71623	H.sapiens ZNF74-1 mRNA > :: gb G27154 G271 54 human STS SHGC-31580.	4e-012	113669	IIII ALU CLASS D WARNING ENTRY IIII	4.1	
2910	U43626	Human chromosome 15q11-q13 putative DNA replication origin in the g- aminobutyric acid receptor b3 and a5 gene cluster	7e-007	2394501	(AF024503) No definition line found [Caenorhabditis elegans]	9.6	
2911	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.3	
2912	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	2688749	(AE001179) conserved hypothetical protein [Borrelia burgdorferi]	2.3	
2913	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.9	
2914	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4	
2915	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
2916	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.004	1209842	(U45423) minus strand repeat motif-containing gene	0.092	

	Ŋ	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2917	X80283	P.polycephalum genomic DNA containing Taq I repetitive element	3.3	<none></none>	<none></none>	<none></none>	
2918	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	<none></none>	<none></none>	<none></none>	
2919	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	<none></none>	<none></none>	<none></none>	
2920	Z97333	Homo sapiens RHCE gene	9e-020	113667	!!!! ALU CLASS B WARNING ENTRY !!!!	4e-005	
2921	AF082350	Homo sapiens bone morphogenetic protein 15 precursor (BMP15) gene, exon 2 and complete cds	1	<none></none>	<none></none>	<none></none>	
2922	L14684	Rattus norvegicus nuclear-encoded mitochondrial elongation factor G mRNA, complete cds.	0	585084	ELONGATION FACTOR G, MITOCHONDRI AL PRECURSOR (MEF-G) >gi 543383 pir S4 0780 translation elongation factor G, mitochondrial - rat >gi 310102	9e-089	
2923	D78335	Human mRNA for 5'-terminal region of UMK, complete cds	e-163	1718058	URIDINE KINASE (URIDINE MONOPHOSPHO KINASE) >gi 471981 (L31783) uridine kinase	7e-072	
2924	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		`	s. Non-Redundant Pro		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2925	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-013	1351922	AMINE OXIDASE PRECURSOR (MONAMINE OXIDASE) (TYRAMINE OXIDASE) >gi 419575 pir B4 1836 amine oxidase (flavin- containing) (EC 1.4.3.4) precursor - Klebsiella pneumoniae >gi 216723 gnl PI D d1001529	5.6	
2926	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds		<none></none>	<none></none>		
2927	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	9.7	
2928	AB018285	Homo sapiens mRNA for KIAA0742 protein, partial cds	0	3882205	(AB018285) KIAA0742 protein [Homo sapiens]	2e-093	
2929	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	3882183	(AB018274) KIAA0731 protein [Homo sapiens]	4e-049	
2930	X94762	H.sapiens DNA for Ki-67 antigen 5'-region (exon 1 & 2)		631020	Kallmann syndrome protein homolog - chicken	5.6	
2931	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	3522948	(AC004411) hypothetical protein [Arabidopsis thaliana]	2e-026	
2932	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.5	

	<u> </u>	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		cds					
2933	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-008	<none></none>	<none></none>	<none></none>	
2934	M18795	Gorilla pseudo- beta- and delta- globin gene intergenic region with 2 Alu repeats.	7e-028	<none></none>	<none></none>	<none></none>	
2935	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0005	<none></none>	<none></none>	<none></none>	
2936	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-007	<none></none>	<none></none>	<none></none>	
2937	U09874	Mus musculus SKD3 mRNA, complete cds.	2e-086	2493735	SKD3 PROTEIN SKD3 [Mus musculus]	6e-036	
2938	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
2939	D38417	Mouse mRNA for arylhydrocarbon receptor, complete cds	e-154	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.4	
2940	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	3879062	(Z81576) predicted using Genefinder	9.2	
2941	AE001368	Plasmodium falciparum chromosome 2, section 5 of 73 of the complete sequence	0.014	<none></none>	<none></none>	<none></none>	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2942	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
2943	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
2944	AF083322	Homo sapiens centriole associated protein CEP110 mRNA, complete cds	e-133	3435244	(AF083322) centriole associated protein CEP110 [Homo sapiens]	9e-015	
2945	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.014	<none></none>	<none></none>	<none></none>	
2946	L07040	pFNeo eukaryotic expression vector, complete sequence.	2e-038	987050	(X65335) lacZ gene product [unidentified cloning vector]	4e-005	
2947	X65319	Cloning vector pCAT-Enhancer	2e-078	987050	(X65335) lacZ gene product [unidentified cloning vector]	1e-013	
2948	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	<none></none>	<none></none>	<none></none>	
2949	AL031844	Human DNA sequence from clone 361H15 on chromosome 22q13.2-13.33, complete sequence [Homo sapiens]	3.2	<none></none>	<none></none>	<none></none>	
2950	AC002186	Homo sapiens (subclone 1_f12 from P1 H115) DNA sequence	2e-037	2072966	(U93570) p40 [Homo sapiens]	4e-013	

	Ŋ	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2951	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-005	4105414	(AF045593) ETS DNA binding protein Yan [Drosophila virilis]	1.4	
2952	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	629692	hypothetical protein - common tobacco tabacum]	4.3	
2953	S60885	LYAR=cell growth regulating nucleolar protein [mice, EL4 cells, mRNA, 1474 nt]	5e-035	2498524	CELL GROWTH REGULATING NUCLEOLAR PROTEIN >gi 423488 pir A4 0683 cell growth regulating nucleolar protein LYAR - mouse >gi 300372 bbs 13 1782	5e-014	
2954	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-012	<none></none>	<none></none>	<none></none>	
2955	Z23090	H.sapiens mRNA for 28 kDa heat shock protein.	1e-063	1709972	60S RIBOSOMAL PROTEIN L10A (CSA-19)	3e-020	
2956	X87817	M.musculus mRNA for Ulip protein	0.0005	<none></none>	<none></none>	<none></none>	
2957	U87997	Enterococcus faecium enterocin B (entB) gene, complete cds		<none></none>	<none></none>	<none></none>	
2958	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.001	<none></none>	<none></none>	<none></none>	
2959	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	<none></none>	<none></none>	<none></none>	

	1	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2960	U95 098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	3e-009	<none></none>	<none></none>	<none></none>	
2961	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	3e-009	<none></none>	<none></none>	<none></none>	
2962	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	<none></none>	<none></none>	<none></none>	
2963	X62025	H.sapiens rod cG-PDE G gene for 3', 5'-cyclic nucleotide phosphodiesteras e	4e-034	728838	!!!! ALU SUBFAMILY SX WARNING ENTRY	9e-006	
2964	AJ223364	Homo sapiens germ-line DNA upstream of Jkappa locus	1e-023	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.2	
2965	Z47046	Human cosmid QLL2C9 from Xq28	3e-020	804808	(M13100) unknown protein [Rattus norvegicus]	7e-005	
2966	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
2967	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	9e-009	464502	PEROXISOMAL TARGETING SIGNAL RECEPTOR (PEROXISOMAL PROTEIN PAS10) (PEROXIN-5) (PTS1 RECEPTOR) >gi 1078412 pir A 49403 tetratricopeptide- repeat protein PAS10 - yeast tetratricopeptide-	9.5	

	N	learest Neighbor		Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					repeat protein [Saccharomyces cerevisiae] >gi 817830 (Z49701) Pas10p [Sa		
2968	AF035940	Homo sapiens MAGOH mRNA, complete cds	3e-050	2306969	(AF007860) xl- Mago [Xenopus laevis]	1e-041	
2969	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
2970	X66297	H.sapiens Alu repeat (terminator 3)	5e-014	<none></none>	<none></none>	<none></none>	
2971	AB007934	Homo sapiens mRNA for KIAA0465 protein, partial cds	0	3413892	(AB007934) KIAA0465 protein [Homo sapiens]	e-118	
2972	X15982	Ascobolus immersus DNA of linear mitochondrial plasmid pAI2 with virus like replication	0.042	<none></none>	<none></none>	<none></none>	
2973	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	<none></none>	<none></none>	<none></none>	

		learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2974	AC002181	Homo sapiens (subclone 2_a12 from BAC H111) DNA sequence	2e-014	3879351	(Z35663) Short region of similarity with glucose-6-phosphate 1-dehydrogenase from Plasmodium falciparum; cDNA EST EMBL:C12945 comes from this gene; cDNA EST yk251d3.3 comes from this gene; cDNA EST yk251d3.5 comes from this	0.69	
2975	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.002	<none></none>	<none></none>	<none></none>	
2976	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	3334221	4- HYDROXYPHEN YLPYRUVATE DIOXYGENASE 4- hydroxyphenylpyr uvate dioxygenase [Mycosphaerella graminicola]	2e-012	
2977	S60885	LYAR=cell growth regulating nucleolar protein	8e-028	2498524	CELL GROWTH REGULATING NUCLEOLAR PROTEIN >gi 423488 pir A4 0683 cell growth regulating nucleolar protein LYAR - mouse >gi 300372 bbs 13 1782	0.72	
2978	U43958	Cloning vector pRcCMV-luc luciferase gene, complete cds	1e-010	335109	(M24873) major structural protein [Rhesus macaque polyomavirus]	1.1	

	Nearest Neighbor			Nearest Neighbor			
200		astN vs. Genbank)	I D V / 4 V / D	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2979	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-014	399294	CYTOCHROME P450 XXIA3 (STEROID 21- HYDROXYLASE) (P450-C21) >gi 2117374 pir A 32525 steroid 21- monooxygenase (EC 1.14.99.10) cytochrome P450 21A1 - pig >gi 164560 (M83939) steroid 21-hydroxylase	3.5	
2980	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	1169449	PROBABLE EARLY E4 33 KD PROTEIN	1.9	
2981	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.2	
2982	Z11711	H.sapiens gene for alpha-2 macroglobulin, exon 1	2e-014	728835	!!!! ALU SUBFAMILY SC WARNING ENTRY	4.2	
2983	M76363	Human (Papua New Guinean) Mitochondrial DNA control region, sequence 130.	1e-053	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.9	
2984	U21228	Promoter-probe vector pCG1408, complete sequence.	3e-049	<none></none>	<none></none>	<none></none>	
2985	X52994	Sheep mRNA for CD3 gamma subunit (partial)	0.005	1084987	cryptogene protein G4 - Sauroleishmania tarentolae (strain LEM125)	2.6	
2986	X52994	Sheep mRNA for CD3 gamma subunit (partial)	0.005	1084987	cryptogene protein G4 - Sauroleishmania tarentolae (strain LEM125)	2.6	
2987	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2988	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
2989	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	123398 3881655	OCTAMER-BINDING TRANSCRIPTIO N FACTOR 1 (OTF-1) (NF-A1) >gi 104811 pir A3 4873 transcription factor Oct-1, octamer-binding- chicken >gi 212467	3.2	
2990	093102	mitotic phosphoprotein 90 mRNA, complete cds	16-010	3881655	(Z82090) similar to Alpha-2-macroglobulin family (3 domains); cDNA EST EMBL:D67694 comes from this gene [Caenorhabditis elegans]	6e-019	
2991	AB018270	Homo sapiens mRNA for KIAA0727 protein, partial cds	0	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.5	
2992	U58745	Caenorhabditis elegans cosmid C10G6.	1.2	2677839	(AF023476) meltrin-L precursor [Homo sapiens]	0.24	
2993	X17051	E.gracilis DNA for ribosomal protein operon	0.13	<none></none>	<none></none>	<none></none>	
2994		D.melanogaster Cpo 61.1 gene for couch potato protein.	1.1		(Y12781) transducin (beta) like 1 protein [Homo sapiens]	6e-017	
2995		Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	8e-008		PARA- AMINOBENZOA TE SYNTHASE Streptomyces griseus > gi 388263 (M93058) p- aminobenzoic acid synthase [Streptomyces griseus]	4.2	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
2996	U11270	Human antithrombin III gene, exon 1 and partial cds.	9e-020	113670	!!!! ALU CLASS E WARNING ENTRY !!!!	0.16	
2997	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	3024528	RAS-RELATED PROTEIN RAB2BV	1.1	
2998	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.17	
2999	U51670	Barbus barbus x Barbus meridionalis microsatellite clone no.77	0.13	<none></none>	<none></none>	<none></none>	
3000	U79776	Mus musculus ajuba (Ajuba) mRNA, complete cds	4e-094	1710382	(U79776) ajuba; jub [Mus musculus]	8e-037	
3001	U79776	Mus musculus ajuba (Ajuba) mRNA, complete cds	4e-094	1710382	(U79776) ajuba; jub [Mus musculus]	8e-037	
3002	U79776	Mus musculus ajuba (Ajuba) mRNA, complete cds	e-100	1710382	(U79776) ajuba; jub [Mus musculus]	8e-019	
3003	U79776	Mus musculus ajuba (Ajuba) mRNA, complete cds	e-100	1710382	(U79776) ajuba; jub [Mus musculus]	8e-019	
3004	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0005	482227	hypothetical protein T07C4.9 - Caenorhabditis elegans >gi 3879509 gnl PI D e1349070 (Z29443) similar to Annexin; cDNA EST EMBL:C10640 comes from this gene; cDNA EST EMBL:C12433 comes from this	0.64	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					gene; cDNA EST yk192f7.5 comes from this gene; cDNA EST yk318c1		
3005	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	1944590	(Z94121) hypothetical protein Rv3899c	7.8	
3006	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
3007	U40603	Rattus norvegicus rat-slowpoke- alpha mRNA, complete cds	0.12	1082665	oligodendrocyte- specific proline- rich protein 2 - human >gi 1408050 gnl PI D d1006205 (D28114) MOBP [Homo sapiens]	0.22	
3008	AF044081	Rattus norvegicus steroidogenic acute regulatory protein (StAR) mRNA, complete cds	1.1	2213519	(Z97050) sigG [Mycobacterium tuberculosis]	3.1	
3009	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
3010	X13345	Human gene for plasminogen activator inhibitor 1	1e-009	<none></none>	<none></none>	<none></none>	
3011	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	395338	(X66924) helix- loop-helix protein [Homo sapiens]	0.85	

	1	Nearest Neighbor		Nearest Neighbor			
	(B)	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3012	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
3013	D78335	Human mRNA for 5'-terminal region of UMK, complete cds	e-101	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.2	
3014	U03887	Human BXP20 gene.	6e-005	<none></none>	<none></none>	<none></none>	
3015	U43194	Mus musculus rhophilin mRNA, complete cds	4e-044	1176422	(U43194) rhophilin [Mus musculus]	7e-020	
3016	AC004507	Homo sapiens chromosome 5, P1 clone 798F12 (LBNL H82), complete sequence [Homo sapiens]	1.2	<none></none>	<none></none>	<none></none>	
3017	X63436	B.taurus mRNA for poly(A) polymerase	0	464345	POLY(A) POLYMERASE (PAP) polynucleotide adenylyltransferas e [Bos taurus]	6e-065	
3018	M98512	Human NFG genomic fragment.	1e-021	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.095	
3019	AJ005016	Homo sapiens mRNA for putative ABC transporter, partial	e-159	3005931	(AJ005016) ABC transporter [Homo sapiens]	2e-039	
3020	AJ006778	Homo sapiens mRNA for DRIM protein	1e-053	<none></none>	<none></none>	<none></none>	
3021	X65319	Cloning vector pCAT-Enhancer	3e-081	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	
3022	U14698	Human Alu-Sb2 repeat, clone HSB-8P.	1e-040	728834	!!!! ALU SUBFAMILY SB2 WARNING ENTRY	0.0001	

	Nearest Neighbor			Nearest Neighbor			
QT-C		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3023	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	3218396	(AL023860) hypothetical protein	0.0003	
3024	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1.20E-01	<none></none>	<none></none>	<none></none>	
3025	Z59351	H.sapiens CpG DNA, clone 151a12, reverse read cpg151a12.rt1a.	3e-020	1079063	deep orange protein - fruit fly (Drosophila melanogaster) >gi 798832 (X86683) deep orange (dor)	9.90E-02	
3026	AB014564	Homo sapiens mRNA for KIAA0664 protein, partial cds	e-164	2498095	5E5 ANTIGEN >gi 1085558 pir J C4163 DNA- binding protein 5E5 - rat norvegicus] >gi 1581020 prf 2 116328A DNA- binding protein 5E5 [Rattus norvegicus]	3.2	
3027	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	<none></none>	<none></none>	<none></none>	
3028	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	<none></none>	<none></none>	<none></none>	
3029	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	<none></none>	<none></none>	<none></none>	
3030		Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4.00E-12	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	

	Nearest Neighbor			Nearest Neighbor			
250	,	lastN vs. Genbank)	I w	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3031	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1.20E-01	<none></none>	<none></none>	<none></none>	
3032	AF070523	Homo sapiens JWA protein mRNA, complete cds	0.00E+00	<none></none>	<none></none>	<none></none>	
3033	Z19055	B.aphidicola tryptophan operon	0.41	<none></none>	<none></none>	<none></none>	
3034	Z19055	B.aphidicola tryptophan operon	0.41	<none></none>	<none></none>	<none></none>	
3035	Z19055	B.aphidicola tryptophan operon	0.41	<none></none>	<none></none>	<none></none>	
3036	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	7.00E-07	<none></none>	<none></none>	<none></none>	
3037	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
3038	AF064482	Homo sapiens natural resistance- associated macrophage protein 2 (NRAMP2) gene, exons 16 and 16A, alternatively spliced IRE form, complete cds	0	<none></none>	<none></none>	<none></none>	
3039	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1.20E-01	<none></none>	<none></none>	<none></none>	
3041	U28153	Caenorhabditis elegans UNC-76 (unc-76) gene, complete cds.	0.38	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3042	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5.00E-03	1079063	deep orange protein - fruit fly (Drosophila melanogaster) >gi 798832 (X86683) deep orange (dor)	0.23	
3043	Û95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5.00E-03	1079063	deep orange protein - fruit fly (Drosophila melanogaster) >gi 798832 (X86683) deep orange (dor)	0.23	
3044	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	2.00E-05	<none></none>	<none></none>	<none></none>	
3045	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
3046	U63810	Homo sapiens WD40 protein Ciao 1 mRNA, complete cds	0.00E+00	3219331	(AC004020) Unknown gene product [Homo sapiens]	2e-097	
3047	M21533	Human MHC class I lymphocyte antigen (HLA-E)	2e-005	120467	V-FOS/FOX TRANSFORMIN G PROTEIN murine osteosarcoma virus (provirus) (fragment)	9.9	
3048	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	6e-006	462702	NEUROFILAME NT TRIPLET H PROTEIN (200 KD NEUROFILAME NT PROTEIN) (NF-H)	2.6	
3049	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3050	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.6	
3051	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5.00E-04	3116127	(AL023287) hypothetical protein	6.9	
3052	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7.00E-06	586875	HYPOTHETICAL 29.2 KD PROTEIN IN METS-KSGA INTERGENIC REGION >gi 2127033 pir S 66068 hypothetical protein - Bacillus subtilis subtilis] >gi 2632306 gnl PI D e1181972 (Z99104) similar to hypothetical proteins [Bacillus subtilis]		
3053	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	1326350	(U58748) similar to potential transmembrane domains in S. cerevisiae nulcear division RFT1 protein (SP:P38206)	0.035	
3054	Y10938	Homo sapiens retroviral-like sequence S71, 5LTR and env- like sequence	6e-016	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.4	
3055	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-05	<none></none>	<none></none>	<none></none>	
3056	AE000723	Aquifex aeolicus section 55 of 109 of the complete genome	1.20E+00	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3057	U18055	Lycopersicon esculentum 1- aminocyclopropa ne-1-carboxylate synthase (LE- ACS3) DNA, partial cds	1.10E+00	<none></none>	<none></none>	<none></none>	
3058	AJ006025	Cicer arietinum mRNA for acyl- coA synthetase, partial	0.38	<none></none>	<none></none>	<none></none>	
3059	AJ006025	Cicer arietinum mRNA for acyl- coA synthetase, partial	0.38	<none></none>	<none></none>	<none></none>	
3060	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.014	<none></none>	<none></none>	<none></none>	
3061	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.014	<none></none>	<none></none>	<none></none>	
3062	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3.00E-10	3880303	[Caenorhabditis elegans]	4.10E-02	
3063	AE000723	Aquifex aeolicus section 55 of 109 of the complete genome	1.20E+00	<none></none>	<none></none>	<none></none>	
3064	Y14352	Gallus gallus gene encoding neurofascin, exons 31 & 31	0.042	995644	(Z54206) UL38 [Bovine herpesvirus 1] >gi 1149580 (Z49078) UL38 [Bovine herpesvirus 1] >gi 2653309 gnl PI D e1187305	1.9	
3065	AE000723	Aquifex aeolicus section 55 of 109 of the complete genome	1.20E+00	<none></none>	<none></none>	<none></none>	

	ľ	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3066	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2.00E-03	<none></none>	<none></none>	<none></none>	
3067	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2.00E-03	<none></none>	<none></none>	<none></none>	
3068	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2.00E-03	<none></none>	<none></none>	<none></none>	
3069	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.38	1395143	(D86080) aniline dioxygenase reductase component [Acinetobacter sp.] dioxygenase reductase component [Acinetobacter sp.]	9.00E-05	
3070	AE001398	Plasmodium falciparum chromosome 2, section 35 of 73 of the complete sequence	0.0005	<none></none>	<none></none>	<none></none>	
3071	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.014	<none></none>	<none></none>	<none></none>	
3072	D16902	Human HepG2 3' region cDNA, clone hmd2h10	2.00E-49	<none></none>	<none></none>	<none></none>	
3073	Z26494	S.cerevisiae genes for histone H2A and H2B, trehalase, and hexaprenyl pyrophosphate synthetase	1.1	3581891	(AL031540) hypothetical wtf3 protein	9.70E+00	

	N	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3074	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2.00E-05	2224921	(AF000606) insect intestinal mucin IIM22 [Trichoplusia ni]	1e-005	
3075	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.37	<none></none>	<none></none>	<none></none>	
3076	U18157	Human HLA class I genomic survey sequence.	2.00E-05	<none></none>	<none></none>	<none></none>	
3077	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4.20E-02	2622750	(AE000921) DNA topoisomerase I [Methanobacteriu m thermoautotrophic um]	2.5	
3078	AF022789	Homo sapiens ubiquitin hydrolyzing enzyme I	0.00E+00	<none></none>	<none></none>	<none></none>	
3079	U18055	Lycopersicon esculentum 1- aminocyclopropa ne-1-carboxylate synthase (LE- ACS3) DNA, partial cds	1.10E+00	<none></none>	<none></none>	<none></none>	
3080	AF022388	Caenorhabditis elegans putative transcription factor MAB-3 (mab-3) gene, complete cds	1.40E-02	3747107	(AF095741) unknown [Rattus norvegicus]	6e-012	
3081	AF084594	Plasmodium falciparum erythrocyte membrane protein 1 type w (var) gene, partial cds	1.20E+00	3132802	(AF063223) pol protein [Human immunodeficiency virus type 1]	1.2	
3082	D16902	Human HepG2 3' region cDNA, clone hmd2h10	2.00E-49	<none></none>	<none></none>	<none></none>	
3083	X65709	A.carrageenovora gene for arylsulfatase	0.014	<none></none>	<none></none>	<none></none>	

		learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3084	AF060246	Mus musculus strain C57BL/6 zinc finger protein 106 (Zfp106) mRNA,	2e-078	3372657	(AF060246) zinc finger protein 106 [Mus musculus]	1e-031	
		H3a-a allele, complete cds					
3085	AF037332	Homo sapiens Eph-like receptor tyrosine kinase hEphB1b (EphB1) mRNA, complete cds	3.70E-01	<none></none>	<none></none>	<none></none>	
3086	U17579	Human growth hormone- releasing hormone receptor gene, alternatively spliced forms a, b, and c, partial cds	0.053	<none></none>	<none></none>	<none></none>	
3087	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.39	2950453	(AL022071) beta- transducin	2.00E-05	
3088	U67479	Methanococcus jannaschii section 21 of 150 of the complete genome	0.005	<none></none>	<none></none>	<none></none>	
3089	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-010	3283350	(AF062378) calmodulin- binding protein SHA1 [Mus musculus]	3e-006	
3090	Z59351	H.sapiens CpG DNA, clone 151a12, reverse read cpg151a12.rt1a.	3e-020	1079063	deep orange protein - fruit fly (Drosophila melanogaster) >gi 798832 (X86683) deep orange (dor)	9.90E-02	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
3091	U95102	Xenopus laevis	2e-005	1125753	(U42833) coded	1.00E-17	
		mitotic			for by C. elegans		
		phosphoprotein			cDNA		
		90 mRNA,			CEESN37F;		
		complete cds			Similar to		
					ammonium		
					transport protein.		
					[Caenorhabditis		
					elegans]	0 000	
3092	AF021834	Homo sapiens	e-172	125932	TISSUE FACTOR	9e-032	
		tissue factor			PATHWAY		
		pathway inhibitor			INHIBITOR PRECURSOR		
		beta (TFPIbeta)			(TFPI)		
		mRNA, complete			(LIPOPROTEIN-		
		cus			ASSOCIATED		
					COAGULATION		
					INHIBITOR)		
					(LACI)		
					(EXTRINSIC		
					PATHWAY		
					INHIBITOR)		
					(EPI) precursor -		
					human >gi 180546		
					(J03225)		
					lipoprotein- associated		
					coagulation		
					inhibitor precursor		
					associated		
					coagulation		
3093	AJ006778	Homo sapiens	0.00E+00	3242214	(AJ006778) DRIM	3e-095	
	210000770	mRNA for DRIM	3.332		protein [Homo		
		protein			sapiens]		
3094	AJ006778	Homo sapiens	0.00E+00	3242214	(AJ006778) DRIM	3e-095	
		mRNA for DRIM			protein [Homo		
		protein			sapiens]		
3095	U95098	Xenopus laevis	6e-005	<none></none>	<none></none>	<none></none>	
		mitotic					
		phosphoprotein					
	ı	44 mRNA, partial					
	4 700 (====	cds		20.4001.4	(A 100/330) DDT (9.005.02	
3096	AJ006778	Homo sapiens	0	3242214	(AJ006778) DRIM	8.00E-93	
		mRNA for DRIM			protein [Homo		
2005	1105004	protein	20.005	<none></none>	sapiens] <none></none>	<none></none>	
3097	U95094	Xenopus laevis XL-INCENP	2e-005	<nune></nune>	\NUNE>	-NONE>	
		(XL-INCENP)					
		mRNA, complete					
		mid va, complete		<u></u>			

		Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)	D 111111	(BlastX vs. Non-Redundant Proteins) ACCESSION DESCRIPTION P VALUE				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	PVALUE		
עו		cds						
		cus						
3098	U95094	Xenopus laevis	3.00E-09	1850115	(Z86089) fadD2	1.4		
		XL-INCENP			[Mycobacterium			
		(XL-INCENP)			tuberculosis]			
		mRNA, complete						
2000	1105004	cds	1e-009	<none></none>	<none></none>	<none></none>		
3099	U95094	Xenopus laevis XL-INCENP	16-009	\NONE/	NONE	NONE		
		(XL-INCENP)						
		mRNA, complete						
		cds						
3100	U95094	Xenopus laevis	1e-009	<none></none>	<none></none>	<none></none>		
		XL-INCENP						
		(XL-INCENP)						
		mRNA, complete						
2101	1167006	cds Bacillus	1.1	2102696	(U72761)	1.90E+00		
3101	U67986	megaterium	1.1	2102090	karyopherin beta 3	1.901		
		anthranilate			[Homo sapiens]			
		synthase (trpD)			[
		gene, partial cds,						
		indole glycerol						
		phosphate						
		synthetase N-						
1		phosphoribosylan						
1		thranilate isomerase (trpF)						
İ		gene partial cds						
3102	U95102	Xenopus laevis	3e-010	2072296	(U95098) mitotic	5.5		
13.02	0,3102	mitotic			phosphoprotein 44			
		phosphoprotein			[Xenopus laevis]			
		90 mRNA,						
		complete cds						

	ľ	Nearest Neighbor			Nearest Neighbor			
	,	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3103	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1.00E-10	135554	TETRACYCLINE RESISTANCE PROTEIN Bacillus cereus plasmid pBC16 >gi 72838 pir YTS OG tetracycline resistance protein - Streptococcus agalactiae plasmid pMV158 >gi 80428 pir JQ1 211 tetracycline resistance protein - Bacillus sp. plasmid pTB19 >gi 151696 (M63	1.4		
3104	AJ006778	Homo sapiens mRNA for DRIM protein	0	3242214	(AJ006778) DRIM protein [Homo sapiens]	8.00E-93		
3105	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	<none></none>	<none></none>	<none></none>		
3106	M60562	Mus musculus Mhc class II A beta polypeptide, partial cds (exons 3 and 4)	1.10E+00	<none></none>	<none></none>	<none></none>		
3107	-U91985	Human DNA fragmentation factor-45 mRNA, complete cds	e-133	2810997	DNA FRAGMENTATI ON FACTOR-45 factor-45 [Homo sapiens]	7e-013		
3108	¥11455	S.salar microsatellite DNA, CA-repeat (AC)11.5	3.50E-01	3879530	(Z49130) cDNA EST yk486b9.3 comes from this gene; cDNA EST yk486b9.5 comes from this gene	0.0001		
3109	¥11455	S.salar microsatellite DNA, CA-repeat (AC)11.5	3.50E-01	3879530	(Z49130) cDNA EST yk486b9.3 comes from this gene; cDNA EST yk486b9.5 comes from this gene	0.0001		

	N	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
3110	AF052135	Homo sapiens	4e-033	4098124	(U73522) STAM	5e-033	
		clone 23625			SH3 domain		
		mRNA sequence			associating		
					molecule [Homo		
					sapiens]	210215	
3111	U95098	Xenopus laevis	6e-005	<none></none>	<none></none>	<none></none>	
		mitotic phosphoprotein					
		44 mRNA, partial					
		cds					
3112	U95102	Xenopus laevis	9e-009	1351538	HYPOTHETICAL	1.4	
	0,0,0,0	mitotic			PROTEIN MG306		
		phosphoprotein			Mycoplasma		
	:	90 mRNA,			genitalium (SGC3)		
		complete cds			>gi 3844885		
					(U39711)		
					conserved		
					hypothetical		
					protein [Mycoplasma		
					genitalium]		
3113	L78777	Homo sapiens	1.30E-01	<none></none>	<none></none>	<none></none>	
3113	E/G///	(subclone 2_b8	1.502 01	110112	1,01,2	1.01.12	
		from P1 H49)					
		DNA sequence					
3114	U95102	Xenopus laevis	0.002	<none></none>	<none></none>	<none></none>	
		mitotic					
		phosphoprotein					
		90 mRNA, complete cds					
3115	U29917	Human AMP	3.00E-10	<none></none>	<none></none>	<none></none>	
3.13	02//1/	deaminase	3.002 10	4.02.12	2101115	210112	
		(AMPD3) gene,					
		exon 8 and 9.					
3116	U95102	Xenopus laevis	9e-009	<none></none>	<none></none>	<none></none>	
		mitotic					
		phosphoprotein					
		90 mRNA,					
2117	A E001029	complete cds	0.14	/NONE>	AIONIES	∠NONE>	
3117	AE001038	Archaeoglobus fulgidus section	U.14	<none></none>	<none></none>	<none></none>	
		69 of 172 of the					
		complete genome					
3118	AF042378	Homo sapiens	0	2801699	(AF042378)	4e-080	
	1 1 0 1 2 0 7 0	spindle pole body	ŭ		spindle pole body	550	
		protein spc98			protein spc98		
		homolog GCP3			homolog GCP3		
		mRNA, complete					
		cds					
			207				

	Nearest Neighbor			Nearest Neighbor			
18	,	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3119	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
3120	Ü95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	1351538	HYPOTHETICAL PROTEIN MG306 Mycoplasma genitalium (SGC3) >gi 3844885 (U39711) conserved hypothetical protein [Mycoplasma genitalium]	1.4	
3121	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7.00E-07	133361	DNA-DIRECTED RNA POLYMERASE III 128 KD POLYPEPTIDE (RNA POLYMERASE III SUBUNIT 2) 2.7.7.6) III second-largest chain - fruit fly polymerase III second-largest subunit [Drosophila melanogaster]	4.40E+00	
3122	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2.00E-05	<none></none>	<none></none>	<none></none>	
3123	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2.00E-05	<none></none>	<none></none>	<none></none>	
3124	AJ011981	Homo sapiens mRNA sequence, IMAGE clone 417820	2.00E-69	461950	DPY-19 PROTEIN elegans >gi 156300 (L12018) putative [Caenorhabditis elegans]	2e-026	

	, l	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3125	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
3126	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.60E+00	
3127	M26216	Rat (lambda 20BH0.1) L-type 6-phosphofructo- 2-kinase/fructose- 2, 6- bisphosphatase	4.10E-02	205752	(M94288) Nopp140 [Rattus norvegicus]	1.1	
3128	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5.00E-03	<none></none>	<none></none>	<none></none>	
3129	<none></none>	<none></none>	<none></none>	730888	OCTAPEPTIDE- REPEAT PROTEIN T2 >gi 296382	5.2	
3130	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5.00E-03	<none></none>	<none></none>	<none></none>	
3131	X65446	H.sapiens gene locus DXS278 (S232-RU2) DNA	6e-011	119110	EBNA-1 NUCLEAR PROTEIN herpesvirus 4 (strain B95-8) >gi 1334880 (V01555) BKRF1 encodes EBNA-1 protein, latent cycle gene. [Human herpesvirus 4]	1e-005	

		Nearest Neighbor		Nearest Neighbor			
O.D.O.		lastN vs. Genbank)	DAVALIE	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3132	X65446	H.sapiens gene locus DXS278 (S232-RU2) DNA	6e-011	119110	EBNA-1 NUCLEAR PROTEIN herpesvirus 4 (strain B95-8) >gi 1334880 (V01555) BKRF1 encodes EBNA-1 protein, latent cycle gene. [Human herpesvirus 4]	1e-005	
3133	X72219	C.pasteurianum gap gene	0.015	<none></none>	<none></none>	<none></none>	
3134	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
3135	Z26494	S.cerevisiae genes for histone H2A and H2B, trehalase, and hexaprenyl pyrophosphate synthetase	1.1	3581891	(AL031540) hypothetical wtf3 protein	9.70E+00	
3136	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.60E+00	
3137	AL010234	Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS *** from contig 4-55, complete sequence	0.37	1213606	(X95910) ftsA [Campylobacter jejuni]	4.2	
3138	U28153	Caenorhabditis elegans UNC-76 (unc-76) gene, complete cds.	0.39	<none></none>	<none></none>	<none></none>	
3139	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8.00E-07	<none></none>	<none></none>	<none></none>	

100	l l	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3140	J05073	Human phosphoglycerate mutase (PGAM- M) gene, complete cds.	1.00E-13	281501	phenylalanine tRNA ligase (EC 6.1.1.20) beta chain - Thermus aquaticus	7		
3141	M90656	Human gamma- glutamylcysteine synthetase (GCS) mRNA, complete cds.	0	1346190	GLUTAMATE CYSTEINE LIGASE CATALYTIC SUBUNIT (GAMMA- GLUTAMYLCYS TEINE SYNTHETASE) glutamate cysteine ligase (EC 6.3.2.2) heavy chain - human >gi 183039 (M90656) gamma- glutamylcysteine synthetase [Homo sapiens]	2.00E-71		
3142	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	8e-006	951325	(U31517) nuclear receptor XR78E/F [Drosophila melanogaster]	9.4		
3143	AF053551	Homo sapiens metaxin 2 (MTX2) mRNA, nuclear gene encoding mitochondrial protein, complete cds	0.00E+00	3283049	(AF053551) metaxin 2 [Homo sapiens]	1.00E-79		
3144	AF088034	Homo sapiens full length insert cDNA clone ZC24F03	e-125	1353059	HYPOTHETICAL 27.4 KD PROTEIN IN MER2-BNA1 INTERGENIC REGION >gi 1077874 pir S 57042 hypothetical protein YJR024c - yeast (Saccharomyces cerevisiae)	9e-023		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					>gi 1015663 (Z49524) ORF YJR024c gene product [Saccharomyces cerevisiae]		
3145	AF087973	Homo sapiens full length insert cDNA clone YU79H10	1e-033	<none></none>	<none></none>	<none></none>	
3146	AF032456	Homo sapiens ubiquitin conjugating enzyme G2	8.00E-07	<none></none>	<none></none>	<none></none>	
3147	Y12259	R.norvegicus mRNA for Kir3.1 protein	6e-058	<none></none>	<none></none>	<none></none>	
3148	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2.00E-07	<none></none>	<none></none>	<none></none>	
3149	X97154	D.willistoni mitochondrial 12S rRNA gene	1.20E+00	3875246	(Z81490) similar to WD domain, Gbeta repeats (2 domains); cDNA EST EMBL:T00482 comes from this gene; cDNA EST EMBL:T00923 comes from this gene; cDNA EST yk449d4.3 comes from this gene; cDNA EST yk449d4.5 comes from this gene; cDNA EST yk449d4.5 comes from this gen	7e-016	
3150	U17247	Saccharomyces cerevisiae chromosome XII cosmid L2142	1.20E-01	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	,	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3151	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	172012	(M12087) thr- tRNA-synthetase [Saccharomyces cerevisiae]	0.21	
3152	L78777	Homo sapiens (subclone 2_b8 from P1 H49) DNA sequence	1.30E-01	<none></none>	<none></none>	<none></none>	
3153	AF053551	Homo sapiens metaxin 2 (MTX2) mRNA, nuclear gene encoding mitochondrial protein, complete cds	0.00E+00	3283049	(AF053551) metaxin 2 [Homo sapiens]	1.00E-79	
3154	X53616	C.domesticus calnexin (pp90) mRNA	1.1	<none></none>	<none></none>	<none></none>	
3155	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.043	<none></none>	<none></none>	<none></none>	
3156	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.002	3327080	(AB014533) KIAA0633 protein [Homo sapiens]	4.2	
3157	U60337	Homo sapiens beta-mannosidase mRNA, complete cds	0	3024091	BETA- MANNOSIDASE PRECURSOR beta-mannosidase [Homo sapiens]	4e-068	
3158	U32790	Haemophilus influenzae Rd section 105 of 163 of the complete genome	1.1	<none></none>	<none></none>	<none></none>	
3159	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.12	<none></none>	<none></none>	<none></none>	

	1	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		•	s. Non-Redundant Pi		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3160	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	1351696	HYPOTHETICAL 30.4 KD PROTEIN C3H1.13 IN CHROMOSOME I >gi 1103514 (Z68144) unknown	1.5	
3161	U50535	Human BRCA2 region, mRNA sequence CG006	4e-012	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	4.5	
3162	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	2132302	hypothetical protein YPR144c - yeast similarity near C-terminus to RNA Polymerase beta subunit (Swiss Prot. accession number P11213) and CCAAT-binding transcription factor (PIR accession number A36368) [Saccharomyces cerevisiae]	4e-022	
3163	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
3164	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	3123086	HYPOTHETICAL PROTEIN MJ1050 Methanococcus jannaschii >gi 1499895 (U67548) conserved hypothetical protein [Methanococcus jannaschii]	1.3	
3165	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.005	<none></none>	<none></none>	<none></none>	
3166	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial	0.005	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor			Nearest Neighbor			
25.0		lastN vs. Genbank)	[B 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
		cds						
3167	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.005	<none></none>	<none></none>	<none></none>		
3168	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	833783	(X14338) NADH:ubiquinone oxidoreductase (428 AA) [Bos taurus]	0.17		
3169	M20918	C.thummi piger haemoglobin (Hb) gene DNA, complete cds.	0.12	2496813	HYPOTHETICAL 59.9 KD PROTEIN B0304.5 IN CHROMOSOME II >gi 1041884 (U39472) B0304.5 gene product [Caenorhabditis elegans]	0.12		
3170	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	100827	NADH dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 4 - wheat mitochondrion	4.1		
3171	U28153	Caenorhabditis elegans UNC-76 (unc-76) gene, complete cds.	0.38	<none></none>	<none></none>	<none></none>		
3172	AJ008065	Chrysolina bankii 16S rRNA gene, mitotype B2	0.045	<none></none>	<none></none>	<none></none>		
3173		Homo sapiens mRNA for KIAA0691 protein, complete cds	7e-057	3327196	(AB014591) KIAA0691 protein [Homo sapiens]	8e-007		
3174		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	3184082	(AL023781) N- terminal acetyltransferase 1 [Schizosaccharom yces pombe]	1e-036		

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3175	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	3219914	HYPOTHETICAL 16.8 KD PROTEIN C30D10.04 IN CHROMOSOME II >gi 2276353 gnl PI D e330328 pombe]	2e-011	
3176	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.002	133361	DNA-DIRECTED RNA POLYMERASE III 128 KD POLYPEPTIDE (RNA POLYMERASE III SUBUNIT 2) 2.7.7.6) III second-largest chain - fruit fly polymerase III second-largest subunit [Drosophila melanogaster]	4.3	
3177	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	2e-006	2429362	(AF020261) proline rich protein [Santalum album]	0.033	
3178	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-007	3641258	(AF064554) ventral anterior homeobox- containing protein 1 [Mus musculus]	0.68	
3179	AB018323	Homo sapiens mRNA for KIAA0780 protein, partial cds	3e-041	3327168	(AB014577) KIAA0677 protein [Homo sapiens]	2e-021	
3180	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	<none></none>	<none></none>	<none></none>	
3181	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	3283350	(AF062378) calmodulin- binding protein SHA1 [Mus musculus]	5e-006	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3182	Z96207	H.sapiens telomeric DNA sequence, clone 12PTEL049, read 12PTELOO049.s eq	8e-008	<none></none>	<none></none>	<none></none>	
3183	AB017026	Mus musculus mRNA for oxysterol-binding protein, complete cds	0	3882265	(AB018315) KIAA0772 protein [Homo sapiens]	2e-091	
3184	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	<none></none>	<none></none>	<none></none>	
3185	X05283	Drosophila melanogaster PKCG7 gene exons 7-14 for protein kinase C	4.6	<none></none>	<none></none>	<none></none>	
3186	AF026069	Homo sapiens phosphomevalona te kinase (HUMPMKI) gene, partial cds	0.42	<none></none>	<none></none>	<none></none>	
3187	AF052573	Homo sapiens DNA polymerase eta (POLH) mRNA, complete cds	0	3510695	(AF052573) DNA polymerase eta [Homo sapiens]	4e-011	
3188	M80198	Human FKBP-12 pseudogene, clone lambda- 512, 5' flank and complete cds.	5e-014	2315521	(AF016452) similar to the beta transducin family	4e-027	
3189	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-008	<none></none>	<none></none>	<none></none>	
3190	AJ001296	Notophthalmus viridescens mRNA for cytokeratin 8	0.38	1175412	HYPOTHETICAL 24.2 KD PROTEIN C13A11.03 IN CHROMOSOME I >gi 984224 (Z54096) unknown	2e-020	

	1	Nearest Neighbor			Nearest Neighbor			
OF O		lastN vs. Genbank)	I S	(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3191	Z60048	H.sapiens CpG DNA, clone 187a9, reverse read cpg187a9.rt1a.	4e-054	547662	HEPATOCYTE NUCLEAR FACTOR 3-BETA HNF-3 beta - mouse >gi 402191 (X74937) HNF- 3beta [Mus musculus]	1e-020		
3192	U95760	Drosophila melanogaster strawberry notch (sno) mRNA, complete cds	3e-071	2078282	(U95760) Sno [Drosophila melanogaster]	3e-068		
3193	L09604	Homo sapiens differentiation- dependent A4 protein mRNA, complete cds.	2e-035	<none></none>	<none></none>	<none></none>		
3194	AF054994	Homo sapiens clone 23832 mRNA sequence	0.12	<none></none>	<none></none>	<none></none>		
3195	AF026069	Homo sapiens phosphomevalona te kinase (HUMPMKI) gene, partial cds	0.42	<none></none>	<none></none>	<none></none>		
3196	AF026069	Homo sapiens phosphomevalona te kinase (HUMPMKI) gene, partial cds	0.42	<none></none>	<none></none>	<none></none>		
3197	AB007918	Homo sapiens mRNA for KIAA0449 protein, partial cds	0.015		GLYCOPROTEIN E PRECURSOR 1 >gi 59566 gnl PID e312380 (X14112) virion glycoprotein E [human herpesvirus 1] >gi 59882 (X02138) glycoprotein gE (Us8) [Human herpesvirus 1] >gi 291496 (L00036) gE protein [Human herpesvirus 1]	8.3		

		Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank))	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE		DESCRIPTION	P VALUE	
3198	L07040	pFNeo eukaryotic expression vector complete sequence.		2072972	(U93572) putative p150 [Homo sapiens]	le-019	
3199	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.002	<none></none>	<none></none>	<none></none>	
3200	M98502	Mus musculus protein encoding twelve zinc finger proteins (pMLZ-4) mRNA, complete cds.	5e-014	<none></none>	<none></none>	<none></none>	
3201	M95098	Bos taurus lysozyme gene (cow 2), complete cds	1.1	3882205	(AB018285) KIAA0742 protein [Homo sapiens]	2e-034	
3202	Ū49169	Dictyostelium discoideum V- ATPase A subunit (vatA) mRNA, complete cds	0.12	2126116	cymH protein - Klebsiella oxytoca >gi 854235	4.2	
3203	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	2911548	(Y15173) E2 protein [Human papillomavirus type 75]	0.39	
3204	Z57610	H.sapiens CpG DNA, clone 187a10, reverse read cpg187a10.rt1a.	7e-090	417134	HEPATOCYTE NUCLEAR FACTOR 3-BETA norvegicus]	5e-019	
3205		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	4104093	(AF031642) urea transporter UT4 [Rattus norvegicus]	0.51	
3206		Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0002	<none></none>	<none></none>	<none></none>	
3207	<none></none>		<none></none>	<none></none>	<none></none>	<none></none>	
3208	<none></none>	<none></none>	<none></none>		(AF006492) FOG [Mus musculus]	3.4	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3209	AF035940	Homo sapiens MAGOH mRNA, complete cds	e-131	2330011	(AF007862) mm- Mago [Mus musculus] >gi 2909828 (AF035939) similar to mago nashi [Mus musculus] >gi 2909830	4e-044	
3210	U49169	Dictyostelium discoideum V- ATPase A subunit (vatA) mRNA, complete cds	0.12	1942101	Porcine Ribonuclease Inhibitor Complexed With Ribonuclease A	1.1	
3211	AF054994	Homo sapiens clone 23832 mRNA sequence	0.12	<none></none>	<none></none>	<none></none>	
3212	AF068627	Mus musculus DNA cytosine-5 methyltransferase 3B2 (Dnmt3b) mRNA, alternatively spliced, complete cds	0.0005	1869835	(Z86099) protein kinase [human herpesvirus 2]	0.86	
3213	X68553	C.elegans repetitive DNA sequence	0.41	854065	(X83413) U88 [Human herpesvirus 6]	7e-007	
3214	X68553	C.elegans repetitive DNA sequence	0.41	854065	(X83413) U88 [Human herpesvirus 6]	7e-007	
3215	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	6e-005	<none></none>	<none></none>	<none></none>	
3216	AF054994	Homo sapiens clone 23832 mRNA sequence	0.12	<none></none>	<none></none>	<none></none>	
3217		Drosophila melanogaster strawberry notch (sno) mRNA, complete cds	3e-071	2078282	(U95760) Sno [Drosophila melanogaster]	3e-068	
3218		P.tetraurelia alpha-51D gene	0.38	<none></none>	<none></none>	<none></none>	

		learest Neighbor		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
410		astN vs. Genbank)	DAZATAT				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION			
3219	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
3220	AF067212	Caenorhabditis elegans cosmid F37F2	0.005	<none></none>	<none></none>	<none></none>	
3221	Y08844	L.esculentum PR1a2 gene	1.1	<none></none>	<none></none>	<none></none>	
3222	Y08844	L.esculentum PR1a2 gene	1.1	<none></none>	<none></none>	<none></none>	
3223	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6.00E-05	<none></none>	<none></none>	<none></none>	
3224	U08214	Rattus sp. DNA binding protein (URE-B1) mRNA, complete cds.	1.1	477513	mesoderm development regulatory protein Sna - mouse >gi 54121 (X67253) sna [Mus musculus]	1.1	
3225	L19713	Human dematin (HRD1) mRNA, complete cds.	0.051	<none></none>	<none></none>	<none></none>	
3226	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.043	2645389	(U83858) NADH dehydrogenase subunit 4 [Onychomys leucogaster]	7.5	
3227	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5.00E-03		(AF034804) LACK [Leishmania major]	3e-011	
3228	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
3229	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.20E+00	

	Nearest Neighbor			Nearest Neighbor			
	,	astN vs. Genbank)		,	(BlastX vs. Non-Redundant Proteins) ACCESSION DESCRIPTION P VALUE		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION			
3230	Ū95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0005	<none></none>	<none></none>	<none></none>	
3231	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0005	<none></none>	<none></none>	<none></none>	
3232	AF036685	Caenorhabditis elegans cosmid C05B10	0.38	<none></none>	<none></none>	<none></none>	
3233	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
3234	AL010153	Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS *** from contig 3-80, complete sequence	6e-005	<none></none>	<none></none>	<none></none>	
3235	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	5.00E-04	<none></none>	<none></none>	<none></none>	
3236	U28153	Caenorhabditis elegans UNC-76 (unc-76) gene, complete cds.	0.39	<none></none>	<none></none>	<none></none>	
3237	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	<none></none>	<none></none>	<none></none>	
3238	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	<none></none>	<none></none>	<none></none>	
3239	X65319	Cloning vector pCAT-Enhancer	5.00E-77	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	

	Ŋ	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3240	AG000140	Homo sapiens genomic DNA, 21q region, clone: T171X2	1.60E-01	2494505	HEPATOCYTE NUCLEAR FACTOR 3 FORKHEAD HOMOLOG 4 (HFH-4) >gi 2137385 pir I4 9734 HNF-3/fork-head homolog-4 - mouse >gi 550488 (L13204) HNF- 3/fork-head homolog-4 [Mus musculus]	7.5	
3241	L77886	Human protein tyrosine phosphatase mRNA, complete cds	1.00E-21	139560	SATELLITE RNA 48 KD PROTEIN	5.9	
3242	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-008	3879988	(Z68318) T21B10.4 [Caenorhabditis elegans]	7.9	
3243	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-007	3184286	(AC004136) hypothetical protein [Arabidopsis thaliana]	7.7	
3244	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	5.00E-04	<none></none>	<none></none>	<none></none>	
3245	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.005	<none></none>	<none></none>	<none></none>	
3246	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.005	<none></none>	<none></none>	<none></none>	
3247	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	2.00E-05	1050849	(X83742) MAP kinase phosphatase [Xenopus laevis]	4.5	

	Nearest Neighbor			Nearest Neighbor			
an a		astN vs. Genbank)	D 1/4 1 1 10	(BlastX vs. Non-Redundant Proteins) ACCESSION DESCRIPTION P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE			P VALUE	
3248	AF084186	Rattus norvegicus alpha-fodrin (A2A) mRNA, complete cds	0.39	3123155	HYPOTHETICAL 49.0 KD TRP- ASP REPEATS CONTAINING PROTEIN F55F8.5 IN CHROMOSOME I family [Caenorhabditis elegans]	5.00E-29	
3249	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5.00E-04	3293508	(AF069188) NADH dehydrogenase 1 [Ephedrus laevicollis]	0.3	
3250	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	3243110	(AF034976) unknown [Pilayella littoralis]	4.6	
3251	M77812	Rabbit myosin heavy chain mRNA, complete cds.	0.58	3876408	(Z81069) Similarity to Yeast hypothetical 65.2 KD protein (SW:P36076); cDNA EST yk393e9.3 comes from this gene; cDNA EST yk393e9.5 comes from this gene [Caenorhabditis elegans]	3.1	
3252	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
3253	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-007	1351841	ISOCITRATE LYASE (ISOCITRASE) lyase [Lycopersicon esculentum]	6.00E+00	
3254	Z50144	R.norvegicus mRNA for kynurenine/alpha- aminoadipate aminotransferase	2.00E-76	1050752	(Z50144) kynurenine/alpha- aminoadipate aminotransferase	6e-033	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
3255	Z50144	R.norvegicus	2.00E-76	1050752	(Z50144)	6e-033	
		mRNA for			kynurenine/alpha-		
		kynurenine/alpha-			aminoadipate		
		aminoadipate			aminotransferase		
		aminotransferase					
3256	U95102	Xenopus laevis	0.043	109340	pepsin (EC	4.5	
		mitotic			3.4.23) II-2/3		
		phosphoprotein			precursor - rabbit		
		90 mRNA,					
2055	V 105100	complete cds	3e-007	3875769	(725((2) similar	4.20E-01	
3257	U95102	Xenopus laevis mitotic	3e-007	38/3/69	(Z35662) similar to Approximately	4.20E-01	
		1			25 cadherin-		
		phosphoprotein 90 mRNA,			repeats, 3 EGF		
		complete cds			domains and one		
		complete cus			Laminin G		
					domain; cDNA		
					EST		
					EMBL:D27303		
					comes from this		
					gene; cDNA EST		
1					EMBL:D27305		
					comes from this		
					gene; cDNA EST		
					EMBL:D27304		
					comes from this		
					gene;		
					>gi 3876224 gnl PI		
2250	AF041059	III	5.90E-02	<none></none>	D e134589 <none></none>	<none></none>	
3258	AF041059	Homo sapiens WSCR4 gene,	3.90E-02	<none></none>	NONE	-NONE>	
		exon 7 and partial					
		cds					
3259	AF054994	Homo sapiens	0.13	<none></none>	<none></none>	<none></none>	
		clone 23832					
		mRNA sequence					
3260	U87266	Arabidopsis	5.60E-01	1175412	HYPOTHETICAL	2e-009	
		thaliana 2,3-			24.2 KD		
		oxidosqualene-			PROTEIN		
		triterpenoid			C13A11.03 IN		
		cyclase mRNA,			CHROMOSOME		
		complete cds			I >gi 984224		
					(Z54096)		
		<u> </u>			unknown		

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3261	AL010240	Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS *** from contig 4-64, complete sequence	1.3	3882205	(AB018285) KIAA0742 protein [Homo sapiens]	5.00E-10	
3262	L20566	Aspergillus niger acid phospatase complete cds.	3.9	3777583	(AF084481) transmembrane protein [Homo sapiens]	5.00E+00	
3263	U12202	Human ribosomal protein S24 (rps24) gene, complete cds	3.80E+00	<none></none>	<none></none>	<none></none>	
3264	U70139	Mus musculus putative CCR4 protein mRNA, partial cds	0	2251234	(U70139) putative CCR4 protein [Mus musculus]	6e-093	
3265	AF055666	Mus musculus kinesin light chain 2 (Klc2) mRNA, complete cds	0.53	3387889	(AF070532) emb- 5 [Homo sapiens]	0.56	
3266	AF077618	Homo sapiens p73 gene, exon 3	0.4	127709	MYOBLAST DETERMINATIO N PROTEIN 1	7.8	
3267	AF072250	Homo sapiens methyl-CpG binding protein MBD4	e-161	3800809	(AF072250) methyl-CpG binding protein MBD4 [Homo sapiens]	2.00E-47	
3268	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	8e-009	886048	(U25686) E93 [Drosophila melanogaster]	1.8	
3269	AG001313	Homo sapiens genomic DNA, 21q region, clone: 125H6N26	0.0005	<none></none>	<none></none>	<none></none>	
3270	U25846	Homarus americanus clone LOB5 farnesoic acid o- methyltransferase mRNA, complete cds.	1.40E-02	<none></none>	<none></none>	<none></none>	

	7	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3271	AF068627	Mus musculus DNA cytosine-5 methyltransferase 3B2 (Dnmt3b) mRNA, alternatively spliced, complete cds	0.0005	1698496	(U53444) LW- amid and MW- amid-containing preprohormone	4.40E+00	
3272	U60022	Mus musculus antigen processing-associated transporter TAP1-k mRNA, complete cds	3.50E+00	2498941	SPLICEOSOME ASSOCIATED PROTEIN 62 spliceosome- associated protein SAP 62 - human >gi 409219	0.23	
3273	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-005	<none></none>	<none></none>	<none></none>	
3274	U24676	Drosophila melanogaster twinstar (tsr) gene, complete cds	1.20E+00	<none></none>	<none></none>	<none></none>	
3275	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1.50E-02	<none></none>	<none></none>	<none></none>	
3276	AF054994	Homo sapiens clone 23832 mRNA sequence	0.13	<none></none>	<none></none>	<none></none>	
3277	AF072250	Homo sapiens methyl-CpG binding protein MBD4	e-161	3800809	(AF072250) methyl-CpG binding protein MBD4 [Homo sapiens]	2.00E-47	
3278	AF054994	Homo sapiens clone 23832 mRNA sequence	0.13	<none></none>	<none></none>	<none></none>	
3279	AF054994	Homo sapiens clone 23832 mRNA sequence	0.13	<none></none>	<none></none>	<none></none>	
3280	AF054994	Homo sapiens clone 23832 mRNA sequence	0.13	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3281	AF054994	Homo sapiens clone 23832 mRNA sequence	0.13	<none></none>	<none></none>	<none></none>	
3282	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	2e-005	<none></none>	<none></none>	<none></none>	
3283	U20281	Gallus gallus clone pNG13 cell division cycle control protein 37 (cdc37) mRNA, complete cds.	0.017	2642625	(AF032118) intersectin [Xenopus laevis]	1.40E+00	
3284	X65279	pWE15 cosmid vector DNA	2e-059	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	
3285	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	<none></none>	<none></none>	<none></none>	
3286	D80005	Human mRNA for KIAA0183 gene, partial cds	0	<none></none>	<none></none>	<none></none>	
3287	U27341	Bos taurus endothelin converting enzyme-2 Sequence 1 from patent US 5736376	1e-096	2136744	endothelin converting enzyme-2 - bovine	2e-047	
3288	M58417	Drosophila melanogaster laminin B2 gene, complete cds.	0.35	1142698	(U26463) NADPH- dependent aldehyde reductase	6.8	
3289	M58417	Drosophila melanogaster laminin B2 gene, complete cds.	0.35	1142698	(U26463) NADPH- dependent aldehyde reductase	6.8	
3290	AF020043	Homo sapiens chromosome-associated polypeptide	0	1785540	(U82626) basement membrane- associated chondroitin proteoglycan Bamacan [Rattus norvegicus]	e-112	

		Nearest Neighbor			Nearest Neighbor			
0700		astN vs. Genbank)	DYAYID		s. Non-Redundant Pr	·		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3291	U57368	Mus musculus EGF repeat transmembrane protein mRNA, complete cds.	O	1336628	(U57368) EGF repeat transmembrane protein [Mus musculus]	e-101		
3292	AB018323	Homo sapiens mRNA for KIAA0780 protein, partial cds	3e-041	3327168	(AB014577) KIAA0677 protein [Homo sapiens]	1e-021		
3293	X65279	pWE15 cosmid vector DNA	2e-059	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015		
3294	X71642	M.musculus GEG-154 mRNA	3e-092	<none></none>	<none></none>	<none></none>		
3295	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	3879362	(Z81113) similar to DnaJ, prokaryotic heat shock protein, Zinc finger, C2H2 type; cDNA EST yk290e12.5 comes from this gene; cDNA EST yk290e12.3 comes from this gene; cDNA EST yk447h4.5 comes from this gene; cDNA EST yk447h4.5 comes from this gene; cDNA EST yk474e4	3e-005		
3296	AB017026	Mus musculus mRNA for oxysterol-binding protein, complete cds	0	3551523	(AB017026) oxysterol-binding protein	e-103		
3297	U43431	Human DNA topoisomerase III mRNA, complete cds.	0	2501242	DNA TOPOISOMERAS E III >gi 1292912	6e-069		
3298	M35296	Human tyrosine kinase arg gene mRNA.	1.1	2135080	epithelial microtubule- associated protein, 115K - human >gi 414115 (X73882) microtubule associated protein	1.8		

		Nearest Neighbor		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	•	astN vs. Genbank)		`			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					[Homo sapiens]		
3299	D50646	Mouse mRNA for SDF2, complete cds	1e-031	2136205	stromal cell- derived factor 2 - human sapiens]	4e-014	
3300	L34732	Homo sapiens T- cell receptor beta (TCRB) mRNA	0.35	3875664	(Z83104) predicted using Genefinder	3e-005	
3301	AF030558	Rattus norvegicus phosphatidylinosi tol 5-phosphate 4- kinase gamma mRNA, complete cds	1e-013	<none></none>	<none></none>	<none></none>	
3302	X03100	Human HLA- SB(DP) alpha gene	2e-018	<none></none>	<none></none>	<none></none>	
3303	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	2950243	(Z98204) extensin [Hordeum vulgare]	2e-005	
3304	Y13631	Clostridium botulinum P-21, P-47 ntnh, bonT genes	1	<none></none>	<none></none>	<none></none>	
3305	Y13631	Clostridium botulinum P-21, P-47 ntnh, bonT genes	1	<none></none>	<none></none>	<none></none>	
3306	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-011	1655830	(U59446) myrosinase- binding protein related protein	0.01	
3307	X17538	Butyrivibrio fibrisolvens end1 gene for endoglucanase	0.12	1001811	(D64005) hypothetical protein	5.2	
3308	D42053	Human mRNA for KIAA0091 gene, complete cds	0	577309	(D42053) KIAA0091 gene product is related to subtilisin. [Homo sapiens]	e-127	

	N	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3309	L81800	Homo sapiens (subclone 2_g9 from P1 H31) DNA sequence	2e-006	<none></none>	<none></none>	<none></none>	
3310	L81800	Homo sapiens (subclone 2_g9 from P1 H31) DNA sequence	2e-006	<none></none>	<none></none>	<none></none>	
3311	K01641	Mouse Ig kappa active V-region from 70Z/3 cells.	3.1	<none></none>	<none></none>	<none></none>	
3312	K01641	Mouse Ig kappa active V-region from 70Z/3 cells.	3.1	<none></none>	<none></none>	<none></none>	
3313	U09954	Human ribosomal protein L9 gene, 5' region and complete cds.	e-114	2136121	ribosomal protein L9 - human >gi 607793	3e-027	
3314	M19735	Homo sapiens beta- hexosaminidase beta chain mRNA, complete cds.	0	179462	(M13519) N- acetyl-beta- glucosaminidase prepro-polypeptide	4e-075	
3315	M31760	Human chromosome 9 t(9;22) breakpoint DNA.	2e-016	2981631	(AB012223) ORF2 [Canis familiaris]	0.018	
3316	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-013	495696	(U00067) C. elegans PAR-3 cell polarity protein	2.5	
3317	U61084	Human phorbolin 3 mRNA, complete cds	0	4097433	(U61084) phorbolin 3 [Homo sapiens]	7e-099	
3318	X95161	H.sapiens brca2 gene exon 11 > :: emb A62786 A62 786 Sequence 27 from Patent WO9719110	5e-024	244126	uroporphyrinogen III synthase, UROIIIS [human, Peptide Mutant, 265 aa]	0.12	
3319	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.9	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3320	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-008	2143490	LGL-1 - mouse >gi 1041889 bbs 1 69033 267 aa] [Mus sp.]	7.2	
3321	U76112	Mus musculus translation repressor NAT1 mRNA, complete cds	1e-013	729818	EUKARYOTIC INITIATION FACTOR 4F SUBUNIT P130 (EIF-4F) (MRNA CAP-BINDING PROTEIN COMPLEX SUBUNIT P130) >gi 539297 pir B4 8086 translation initiation factor eIF-4F TIF4632 - yeast (Saccharomyces cerevisiae) >gi 295677 (L16924) p130 [Saccharomyces cerevisiae	1.9	
3322	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.2	
3323	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-013	495696	(U00067) C. elegans PAR-3 cell polarity protein	2.5	
3324	U43626	Human chromosome 15q11-q13 putative DNA replication origin in the g- aminobutyric acid receptor b3 and a5 gene cluster		2197085	(AF003535) ORF2-like protein [Homo sapiens]	0.0002	
3325	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-010	<none></none>	<none></none>	<none></none>	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3326	Z48561	E.coli perA, perB, perC and perD genes	0.38	2576325	(Y12239) env [porcine endogenous retrovirus]	7.4	
3327	Z48561	E.coli perA, perB, perC and perD genes	0.38	2576325	(Y12239) env [porcine endogenous retrovirus]	7.4	
3328	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	2576325	(Y12239) env [porcine endogenous retrovirus]	7.4	
3329	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	2576325	(Y12239) env [porcine endogenous retrovirus]	7.4	
3330	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1e-010	1362915	protein-tyrosine kinase (EC 2.7.1.112) STK-1 precursor - human	0.5	
3331	X65319	Cloning vector pCAT-Enhancer	3e-081	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	
3332	AB018304	Homo sapiens mRNA for KIAA0761 protein, partial cds	0	3882243	(AB018304) KIAA0761 protein [Homo sapiens]	8e-098	
3333	Y08460	Mus musculus mRNA for Mdes transmembrane protein	1e-085	2225941	(Y08460) Mdes protein [Mus musculus]	8e-071	
3334	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.1	
3335	<none></none>	<none></none>	<none></none>		<none></none>	<none></none>	
3336	·	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	2687928	(AE001118) P115 protein [Borrelia burgdorferi]	5.2	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3337	U94831	Homo sapiens multispanning membrane protein mRNA, complete cds	0	2276460	(U94831) multispanning membrane protein [Homo sapiens]	5e-087	
3338	U14972	Human ribosomal protein S10 mRNA, complete cds.	2e-059	133715	RIBOSOMAL PROTEIN S10	0.0002	
3339	K01254	Human gastrin gene, complete cds.	5e-005	<none></none>	<none></none>	<none></none>	
3340	U08469	Glycine max 3- methylcrotonyl- CoA carboxylase mRNA, biotin- carrier domain, partial cds.	3e-051	3876562	(Z81074) Similarity to Soybean 3- methylcrotonyl- CoA carboxylase (TR:Q42777); cDNA EST EMBL:M75819 comes from this gene; cDNA EST EMBL:M89099 comes from this gene; cDNA EST EMBL:D32737 comes from this gene; cDNA EST EMBL:D32737 comes from this	1e-073	
3341	AB011139	Homo sapiens mRNA for KIAA0567 protein, partial cds	0	3043658	(AB011139) KIAA0567 protein [Homo sapiens]	e-123	
3342	U07615	Rattus norvegicus mucin mRNA, partial cds.	2e-006	2506877	MUCIN 2 PRECURSOR (INTESTINAL MUCIN 2) >gi 454154 (L21998) mucin [Homo sapiens]	0.0007	
3343	AF061749	Homo sapiens tumorous imaginal discs protein Tid56 homolog (TID1) mRNA, complete cds	e-154	3372677	(AF061749) tumorous imaginal discs protein Tid56 homolog	4e-060	

		Nearest Neighbor			Nearest Neighbor			
25.0		lastN vs. Genbank)	I w # 7 4 7 7 7 7	(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3344	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.001	2984587	(AC004472) P1.11659_3 [Homo sapiens]	3e-008		
3345	U45998	Onchocerca volvulus MRS3/MRS4 class mitochondrial solute carrier mRNA, complete cds	2e-008	3880433	(Z66521) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene [Caenorhabditis elegans]	2e-051		
3346	U43626	Human chromosome 15q11-q13 putative DNA replication origin in the g- aminobutyric acid receptor b3 and a5 gene cluster	2e-018	2197085	(AF003535) ORF2-like protein [Homo sapiens]	0.0002		
3347	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.2		
3348	U14972	Human ribosomal protein S10 mRNA, complete cds.	2e-059	133715	40S RIBOSOMAL PROTEIN S10	0.0002		
3349	M80198	Human FKBP-12 pseudogene, clone lambda- 512, 5' flank and complete cds.	1.00E-10	2315521	(AF016452) similar to the beta transducin family	1e-022		
3350	AB011180	Homo sapiens mRNA for KIAA0608 protein, partial cds	5e-077	3043740	(AB011180) KIAA0608 protein [Homo sapiens]	8e-071		
3351	U45858	Zea mays glyceraldehyde-3- phosphate dehydrogenase	4.2	<none></none>	<none></none>	<none></none>		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
3352	U45858	Zea mays	4.2	<none></none>	<none></none>	<none></none>	
1 1		glyceraldehyde-3-					
		phosphate					
3353	AF035940	dehydrogenase Homo sapiens	e-141	2330011	(AF007862) mm-	1e-075	
3333	AF033940	MAGOH mRNA,	6-141	2330011	Mago [Mus	10-075	
		complete cds			musculus]		
		complete cas			>gi 2909828		
					(AF035939)		
					similar to mago		
1					nashi [Mus		
					musculus]		
					>gi 2909830		
3354	AF035940	Homo sapiens	e-141	2330011	(AF007862) mm-	1e-075	
		MAGOH mRNA,			Mago [Mus		
		complete cds			musculus]		
					>gi 2909828 (AF035939)		
					similar to mago		
					nashi [Mus		
					musculus]		
					>gi 2909830		
3355	M24486	Human prolyl 4-	e-147	3876769	(Z69637)	4e-012	
		hydroxylase			Similarity to		
		alpha subunit			Human Prolyl 4-		
		mRNA, complete			hydroxylase alpha subunit		
		cds, clone PA-11.			(SW:P4HA HUM		
					AN); cDNA EST		
					yk219g12.5 comes		
					from this gene;		
					cDNA EST		
					yk319d8.5 comes		
					from this gene;		
					cDNA EST		
					yk339d11.5 comes		
					from this gene; cDNA EST		
		-			yk371c9.3		
3356	Z50144	R.norvegicus	3.00E-93	1050752	(Z50144)	2e-043	
	2001,,	mRNA for			kynurenine/alpha-	=	
		kynurenine/alpha-			aminoadipate	l	
	:	aminoadipate		:	aminotransferase		
		aminotransferase					

		Nearest Neighbor			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	```	astN vs. Genbank)	DYZZIE	•	s. Non-Redundant Production  DESCRIPTION	P VALUE		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION				
3357	M24486	Human prolyl 4- hydroxylase alpha subunit mRNA, complete cds, clone PA-11.	e-147	3876769	(Z69637) Similarity to Human Prolyl 4- hydroxylase alpha subunit (SW:P4HA_HUM AN); cDNA EST yk219g12.5 comes from this gene; cDNA EST yk319d8.5 comes from this gene; cDNA EST yk339d11.5 comes from this gene; cDNA EST yk339d11.5 comes	4e-012		
3358	U83981	Homo sapiens apoptosis associated protein (GADD34) mRNA, complete cds	0	3258618	(U83981) apoptosis associated protein [Homo sapiens]	8.00E-24		
3359	U30817	Bos taurus very- long-chain acyl- CoA dehydrogenase mRNA, nuclear gene encoding mitochondrial protein, complete cds.	1e-010	2765125	(Y11770) very- long-chain acyl- CoA dehydrogenase [Mus musculus]	4e-013		
3360	Z35094	H.sapiens mRNA for SURF-2	5e-097	2498974	SURFEIT LOCUS PROTEIN 2	4e-046		
3361	Z35094	H.sapiens mRNA for SURF-2	5e-097	2498974	SURFEIT LOCUS PROTEIN 2	4e-046		
3362	Z35094	H.sapiens mRNA for SURF-2	5e-097	2498974	SURFEIT LOCUS PROTEIN 2	4e-046		
3363	Z63829	H.sapiens CpG DNA, clone 90h2, forward read cpg90h2.ft1a		1050411	(L43146) nuclear factor I-B1 [Xenopus laevis]	5.4		
3364	AF052573	Homo sapiens DNA polymerase eta (POLH) mRNA, complete cds	0	3510695	(AF052573) DNA polymerase eta [Homo sapiens]	4e-011		

	N	learest Neighbor		Nearest Neighbor				
	(BI	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
ID	ACCESSION	DESCRIPTION	1 VIIDOD	11002031011				
3365	AF092564	Homo sapiens	0	3851586	(AF092564)	6e-052		
		chromosome-			chromosome-			
		associated			associated protein-			
		protein-C			C [Homo sapiens]			
3366	AF031924	Homo sapiens homeobox transcription factor barx2	2.00E-90	<none></none>	<none></none>	<none></none>		
3367	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	419712	probable transposase (insertion sequence IS1138) - Mycoplasma pulmonis (SGC3)	2.6		
3368	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	419712	probable transposase (insertion sequence IS1138) - Mycoplasma pulmonis (SGC3)	2.6		
3369	M24487	Human prolyl 4-hydroxylase alpha subunit mRNA, complete cds, clone PA-15.	e-125	2507090	PROLYL 4- HYDROXYLASE ALPHA SUBUNIT PRECURSOR >gi 66338 pir  DA HUA2 procollagen- proline dioxygenase (EC 1.14.11.2) alpha chain precursor, splice form 2 - human >gi 602675 (U14620) alpha- subunit of prolyl 4-hydroxylase [Homo sapiens]	1e-007		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3370	M24487	Human prolyl 4- hydroxylase alpha subunit mRNA, complete cds, clone PA-15.	e-125	2507090	PROLYL 4- HYDROXYLASE ALPHA SUBUNIT PRECURSOR >gi 66338 pir  DA HUA2 procollagen- proline dioxygenase (EC 1.14.11.2) alpha chain precursor, splice form 2 - human >gi 602675 (U14620) alpha- subunit of prolyl 4-hydroxylase [Homo sapiens]	1e-007	
3371	U45858	Zea mays glyceraldehyde-3- phosphate dehydrogenase	4.2	<none></none>	<none></none>	<none></none>	
3372	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-006	<none></none>	<none></none>	<none></none>	
3373	D50930	Human mRNA for KIAA0140 gene, complete cds	2e-046	<none></none>	<none></none>	<none></none>	
3374	X06461	Herpes simplex virus type I immediate early (IE) gene 3 for transcriptional activator IE175 (= ICP 4)	3.00E-04	2924449	(AL022022) PE_PGRS [Mycobacterium tuberculosis]	4.00E-05	
3375	X06461	Herpes simplex virus type I immediate early (IE) gene 3 for transcriptional activator IE175 (= ICP 4)	3.00E-04	2924449	(AL022022) PE_PGRS [Mycobacterium tuberculosis]	4.00E-05	

		Nearest Neighbor			Nearest Neighbor			
	` .	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)  ACCESSION DESCRIPTION P VALUE				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION			
3376	X85753	Homo sapiens mRNA for CDK8 protein kinase > :: emb A61243 A61 243 Sequence 1 from Patent WO9709432	7e-059	<none></none>	<none></none>	<none></none>		
3377	X76192	Mycoplasma sp. munIM, munIC and munIR genes.	1.2	<none></none>	<none></none>	<none></none>		
3378	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>		
3379	M24486	Human prolyl 4-hydroxylase alpha subunit mRNA, complete cds, clone PA-11.	e-147	3876769	(Z69637) Similarity to Human Prolyl 4- hydroxylase alpha subunit (SW:P4HA_HUM AN); cDNA EST yk219g12.5 comes from this gene; cDNA EST yk319d8.5 comes from this gene; cDNA EST yk339d11.5 comes from this gene; cDNA EST yk339d11.5 comes	4e-012		
3380		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-006	<none></none>	<none></none>	<none></none>		
3381	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	2119163	collagen alpha 1(III) chain precursor - mouse	0.005		
3382	AB009357	Homo sapiens mRNA for TGF- beta activated kinase 1b, complete cds	0	1167506	(D76446) TAK1 (TGF-beta- activated kinase) [Mus musculus]	2e-033		

		learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID			£ 050	14016	(X55654)	1e-014	
3383	D38112	Human mitochondrial DNA, complete sequence	5e-052	14016	cytochrome C oxidase II subunit [Homo sapiens]		
3384	Z96177	H.sapiens telomeric DNA sequence, clone 10QTEL040, read 10QTELOO040.s eq	7e-038	987050	(X65335) lacZ gene product [unidentified cloning vector]	0.035	
3385	Z96177	H.sapiens telomeric DNA sequence, clone 10QTEL040, read 10QTELOO040.s		987050	(X65335) lacZ gene product [unidentified cloning vector]	0.035	
3386	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-011	2384956	(AF022985) No definition line found [Caenorhabditis elegans]	6e-029	
3387	AF010484	Homo sapiens ICI YAC 9IA12, right end sequence		<none></none>	<none></none>	<none></none>	
3388	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	113667	!!!! ALU CLASS B WARNING ENTRY !!!!	0.68	
3389	AJ009761	Homo sapiens mRNA for putative dimethyladenosin e transferase, partial	0	4050050	(AF102147) putative dimethyladenosine transferase [Homo sapiens]	4.00E-46	
3390	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.048	<none></none>	<none></none>	<none></none>	
3391	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.048	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	•	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3392	AL022579	Homo sapiens DNA sequence from clone 47K8 on chromosome Xp11.21-11.23, complete sequence [Homo sapiens]	1e-070	<none></none>	<none></none>	<none></none>	
3393	U37454	Human Down Syndrome region of chromosome 21 genomic sequence, clone A31D6-1H7.	0.12	<none></none>	<none></none>	<none></none>	
3394	AF058954	Homo sapiens GTP-specific succinyl-CoA synthetase beta subunit (SCS) mRNA, partial cds	0	3766199	(AF058954) GTP-specific succinyl-CoA synthetase beta subunit [Homo sapiens]	e-122	
3395	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	3043582	(AB011101) KIAA0529 protein [Homo sapiens]	2e-012	
3396	Z23090	H.sapiens mRNA for 28 kDa heat shock protein.	3e-079	1709972	60S RIBOSOMAL PROTEIN L10A (CSA-19)	2e-025	
3397	D14657	Human mRNA for KIAA0101 gene, complete cds	0	3183216	HYPOTHETICAL PROTEIN KIAA0101 sapiens]	2e-026	
3398	D17577	Mouse mRNA for kinesin-like protein (Kif1b), complete cds		2497524	KINESIN-LIKE PROTEIN KIF1B mouse >gi 407339 gnl PI D d1005029 (D17577) Kif1b [Mus musculus]	1e-048	
3399	AF091078	Homo sapiens clone 559 unknown mRNA, complete sequence	0	4050050	(AF102147) putative dimethyladenosine transferase [Homo sapiens]	1e-048	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID					ALONIE	AIONES	
3400	AC000043	Homo sapiens Chromosome 22q13 Cosmid Clone p74a8, complete sequence [Homo sapiens]	7e-006	<none></none>	<none></none>	<none></none>	
3401	AC000043	Homo sapiens Chromosome 22q13 Cosmid Clone p74a8, complete sequence [Homo sapiens]	7e-006	<none></none>	<none></none>	<none></none>	
3402	AF031924	Homo sapiens homeobox transcription factor barx2	e-156	<none></none>	<none></none>	<none></none>	
3403	AF031924	Homo sapiens homeobox transcription factor barx2	e-157	3882305	(AB018335) KIAA0792 protein [Homo sapiens]	4.5	
3404	L22473	Human Bax alpha mRNA, complete cds.	0	728945	APOPTOSIS REGULATOR BAX, MEMBRANE ISOFORM ALPHA >gi 539664 pir  A4 7538 bcl-2- associated protein bax alpha splice form - human >gi 388166	9e-075	
3405	U04709	Human adenine phosphoribosyltra nsferase (APRT) gene, 3' flanking region		113668	!!!! ALU CLASS C WARNING ENTRY !!!!	0.91	
3406	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-008	3064146	(AF036408) mucin-like protein [Trypanosoma cruzi]	7.6	
3407	AF093268	Rattus norvegicus homer-1c mRNA, complete cds		1913909	(U92079) GLGF- domain protein Homer [Rattus norvegicus]	4e-046	

		Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3408	U47322	Cloning vector DNA, complete sequence.	2.00E-38	987050	(X65335) lacZ gene product [unidentified cloning vector]	3.00E-03	
3409	U78109	Mus musculus prepro-neurturin mRNA, complete cds	1.2	2506998	STANNIOCALCI N (STC) (CORPUSCLES OF STANNIUS PROTEIN) (CS) (HYPOCALCIN) (TELEOCALCIN)	1.2	
3410	Z96177	H.sapiens telomeric DNA sequence, clone 10QTEL040, read 10QTELOO040.s eq	5e-013	<none></none>	<none></none>	<none></none>	
3411	D50930	Human mRNA for KIAA0140 gene, complete cds	0.00E+00	1235974	(X96713) collagen [Globodera pallida]	5.8	
3412	D50930	Human mRNA for KIAA0140 gene, complete cds	2e-046	<none></none>	<none></none>	<none></none>	
3413	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	<none></none>	<none></none>	<none></none>	
3414	L01777	Yersinia pseudotuberculosi s (group IIA) rfb gene cluster	1.20E-01	<none></none>	<none></none>	<none></none>	
3415	D17577	Mouse mRNA for kinesin-like protein (Kif1b), complete cds	e-130	2497524	KINESIN-LIKE PROTEIN KIF1B mouse >gi 407339 gnl PI D d1005029 (D17577) Kif1b [Mus musculus]	1e-049	
3416	AB014597	Homo sapiens mRNA for KIAA0697 protein, partial cds	2e-067	3327208	(AB014597) KIAA0697 protein [Homo sapiens]	6e-050	

	N	learest Neighbor		Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID				2103.45	AIONES	NONES	
3417	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1.00E-11	<none></none>	<none></none>	<none></none>	
3418	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
3419	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
3420	AB014597	Homo sapiens mRNA for KIAA0697 protein, partial cds	2e-067	3327208	(AB014597) KIAA0697 protein [Homo sapiens]	6e-050	
3421	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	
3422	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8.00E-08	1176456	(S79774) bile salt- dependent lipase, BSDL {EC 3.1.1 } [human, fetal pancreas, Peptide Partial, 720 aa] [Homo sapiens]	9.4	
3423	AF100661	Caenorhabditis elegans cosmid H20E11	0.39	<none></none>	<none></none>	<none></none>	
3424	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds		<none></none>	<none></none>	<none></none>	
3425	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9.00E-10	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	•	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3426	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9.00E-10	<none></none>	<none></none>	<none></none>	
3427	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-05	3056592	(AC004255) T1F9.13 [Arabidopsis thaliana]	10	
3428	U89676	Candida albicans putative membrane protein (CSP37) gene, complete cds	0.12	<none></none>	<none></none>	<none></none>	
3429	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	
3430	<none></none>	<none></none>	<none></none>		<none></none>	<none></none>	
3431	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-07	<none></none>	<none></none>	<none></none>	
3432	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds		<none></none>	<none></none>	<none></none>	
3433	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-07	<none></none>	<none></none>	<none></none>	
3434	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4.00E-13		<none></none>	<none></none>	
3435	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4.00E-13	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID			2 065	2227200	(AD014607)	6- 050	
3436	AB014597	Homo sapiens mRNA for KIAA0697 protein, partial cds	2e-067	3327208	(AB014597) KIAA0697 protein [Homo sapiens]	6e-050	
3437	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-06	1360669	collagen alpha 1(V) chain precursor - human sapiens]	1.8	
3438	U65297	Geomys breviceps cytochrome b (cytb) gene, mitochondrial gene encoding mitochondrial protein, complete cds	3.50E+00	<none></none>	<none></none>	<none></none>	
3439	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-06	3914965	TOXIN BMK-X PRECURSOR (BMK10) (BMK M10) (NEUROTOXIN M10) > gi 3138981 (AF062563) neurotoxin M10 precursor [Mesobuthus martensii]	4	
3440	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-06	3914965	TOXIN BMK-X PRECURSOR (BMK10) (BMK M10) (NEUROTOXIN M10) > gi 3138981 (AF062563) neurotoxin M10 precursor [Mesobuthus martensii]	4	
3441	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-011	3413900	(AB007938) KIAA0469 protein [Homo sapiens]	1.40E-02	

		Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3442	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1.00E-11	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.20E+00		
3443	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1.00E-11	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.20E+00		
3444	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8.00E-08	1176456	(\$79774) bile salt- dependent lipase, BSDL {EC 3.1.1 } [human, fetal pancreas, Peptide Partial, 720 aa] [Homo sapiens]	9.4		
3445	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4.00E-13	<none></none>	<none></none>	<none></none>		
3446	U91543	Homo sapiens zinc-finger helicase (hZFH) mRNA, complete cds	1.00E-61	2961557	(AF050199) putative peroxisome microbody protein 175.1	3.70E+00		
3447	X75258	H.sapiens DNA from recombination area	1.40E-02	1143020	(U28974) ORF1 [Spiroplasma virus]	9.5		
3448	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	8.00E-08	<none></none>	<none></none>	<none></none>		
3449	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-07	631089	bat2 protein - human	0.055		

	Nearest Neighbor			Nearest Neighbor			
27.0		astN vs. Genbank)	DVALUE	(BlastX vs. Non-Redundant Proteins)  ACCESSION   DESCRIPTION   P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION		
3450	AL022321	Homo sapiens DNA sequence from PAC 20O8 on chromosome 22q12.1-12.3. Contains exons 13 and 14 of the SLC5A1 (SGLT1) gene for solute carrier family 5 Sodium- Glucose Cot	1.10E+00	3063453	(AC003981) F22O13.15 [Arabidopsis thaliana]	7.2	
3451	AF060798	Homo sapiens myristilated and palmitylated serine-threonine kinase MPSK (MPSK1) mRNA, complete cds	0.00E+00	3372666	(AF060798) myristilated and palmitylated serine-threonine kinase MPSK [Homo sapiens]	2e-067	
3452	AF080399	Drosophila melanogaster mitotic checkpoint control protein kinase BUB1 (Bub1) mRNA, complete cds	1.1	3184082	(AL023781) N- terminal acetyltransferase 1 [Schizosaccharom yces pombe]	1e-033	
3453	AF041259	Homo sapiens breast cancer putative transcription factor (ZABC1) mRNA, complete cds	0.00E+00	3879065	(Z81576) R10E8.3 [Caenorhabditis elegans]	9.7	
3454	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3.70E-01	549359	MINOR CAPSID PROTEIN L2 type 26 >gi 396962 (X74472) late protein [Human papillomavirus type 26]	0.097	
3455	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2.00E-06	2746890	(AF040655) No definition line found [Caenorhabditis elegans]	9.1	

	1	Nearest Neighbor			Nearest Neighbor			
	(B	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3456	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	3874316	(Z81470) predicted using Genefinder	6.8		
3457	V01399	Defective Semliki forest virus RNA. Derived by serial undiluted passaging of the virus in baby hamster kidney cells > :: gb L00017 SFVD IB semliki forest virus defective interfering (18s di) rna di309.	0.98	2496616	HYPOTHETICAL 38.5 KD PROTEIN Y4EE	2.1		
3458	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4.60E-02	<none></none>	<none></none>	<none></none>		
3459	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6.00E-06	<none></none>	<none></none>	<none></none>		
3460	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.014	630844	NADH dehydrogenase chain 2 - fruit fly dehydrogenase subunit 2 [Drosophila erecta]	7.3		
3461	L49035	Gorilla gorilla ABC-transporter (TAP2) mRNA, complete cds	4.70E-01	2058691	(U94836) ERPROT 213-21 [Homo sapiens]	4.3		
3462	U67524	Methanococcus jannaschii section 66 of 150 of the complete genome	4.10E-02		HYPOTHETICAL 82 KD AVIRULENCE PROTEIN IN AVRBS3 REGION >gi 77844 pir  JQ0 317 hypothetical 82K protein - Xanthomonas	7.3		

		Nearest Neighbor		Nearest Neighbor			
	•	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
10					campestris pv. vesicatoria		
3463	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4.00E-12	<none></none>	<none></none>	<none></none>	
3464	U65297	Geomys breviceps cytochrome b (cytb) gene, mitochondrial gene encoding mitochondrial protein, complete cds	3.50E+00	<none></none>	<none></none>	<none></none>	
3465	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
3466	U36266	Human beta- prime-adaptin (BAM22) gene, exons 18 and 19	1.20E+00	<none></none>	<none></none>	<none></none>	
3467	AB018327	Homo sapiens mRNA for KIAA0784 protein, partial cds	0	3882289	(AB018327) KIAA0784 protein [Homo sapiens]	e-103	
3468	AB018327	Homo sapiens mRNA for KIAA0784 protein, partial cds	0	3882289	(AB018327) KIAA0784 protein [Homo sapiens]	e-103	
3469	U667 <b>8</b> 9	Human laminin alpha 2 chain (LAMA2) gene, exon 57	4.80E-02	3873753	(Z66519) similar to phytoene synthase precursor; cDNA EST yk340f7.3 comes from this gene; cDNA EST yk340f7.5 comes from this gene [Caenorhabditis elegans]	3e-006	

		·	Norman Milinkham				
		Nearest Neighbor		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
000		astN vs. Genbank)		`			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3470	AB018327	Homo sapiens mRNA for	9.00E-88	3882289	(AB018327) KIAA0784 protein	9e-022	
		KIAA0784 protein, partial cds			[Homo sapiens]		
3471	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0002	<none></none>	<none></none>	<none></none>	
3472	U67524	Methanococcus jannaschii section 66 of 150 of the complete genome	4.10E-02	140229	HYPOTHETICAL 82 KD AVIRULENCE PROTEIN IN AVRBS3 REGION >gi 77844 pir  JQ0 317 hypothetical 82K protein - Xanthomonas campestris pv. vesicatoria	7.3	
3473	L13972	Homo sapiens beta-galactoside alpha-2,3- sialyltransferase (SIAT4A) mRNA, complete cds	0.005	<none></none>	<none></none>	<none></none>	
3474	L13972	Homo sapiens beta-galactoside alpha-2,3- sialyltransferase (SIAT4A) mRNA, complete cds	0.005	<none></none>	<none></none>	<none></none>	
3475	AL031222	Caenorhabditis elegans cosmid 6R55, complete sequence [Caenorhabditis elegans]	1.10E-01	<none></none>	<none></none>	<none></none>	

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	Nearest Neighbor			Nearest Neighbor			
12.5		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3476	AF070529	Homo sapiens clone 24525 mRNA sequence	0	3879532	(Z49130) cDNA EST EMBL:D74028 comes from this gene; cDNA EST EMBL:D71354 comes from this gene; cDNA EST EMBL:D76320 comes from this gene; cDNA EST yk486c7.3 comes from this gene; cDNA EST yk486c7.5 comes from this gene;	1.50E+00	
3477	U02567	Mus musculus BALB/c T-cell antigen 4-1BB gene, complete cds.	1.30E-01	2414601	(Z99295) phosphatidyl synthase	5e-005	
3478	AB018327	Homo sapiens mRNA for KIAA0784 protein, partial cds	9.00E-88	3882289	(AB018327) KIAA0784 protein [Homo sapiens]	9e-022	
3479	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	2315521	(AF016452) similar to the beta transducin family	2e-006	
3480	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	
3481		H.sapiens mRNA for atr gene > :: gb U76308 HSU7 6308 Human protein kinase ATR mRNA, complete cds > :: emb A61385 A61 385 Sequence 1 from Patent WO9709433	0	1235902	(U49844) FRAP- related protein [Homo sapiens]	3e-051	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3482	Z48633	H.sapiens mRNA for retrotransposon.	e-165	1177607	(X92485) pva l [Plasmodium vivax]	1.9	
3483	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5e-013	111978	mucin - rat	2.6	
3484	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.4	
3485	X77335	A.thaliana gene for methyltransferase	0.13	1401051	(U24160) similar to Dvl-1 product encoded by GenBank Accession Number U10115; dishevelled segment polarity protein homolog [Mus musculus]	3.5	
3486	AF038660	Homo sapiens chromosome 1p33-p34 beta- 1,4- galactosyltransfer ase mRNA, complete cds	е-144	2995442	(Y12510) UDPGal:GlcNAc b1,4 galactosyltransfera se [Homo sapiens]	9e-005	
3487	U65960	Human kinase substrate HASPP28 gene, 5' flanking region and partial cds	1e-021	2120084	reverse transcriptase - mouse >gi 558908	9.7	
3488	AF058907	Homo sapiens pleiotrophin (PTN) gene, exons UV3, UV2 and UV1	8e-060	120806	GAG POLYPROTEIN (CONTAINS: CORE PROTEIN P15; INNER COAT PROTEIN P12; CORE SHELL PROTEIN P30) >gi 74562 pir  FO VDA gag polyprotein - avian spleen necrosis virus (fragment)	5e-005	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					>gi 61758 (V01200) reading frame (gag?) [Spleen necrosis virus]		
3489	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-011	3123086	HYPOTHETICAL PROTEIN MJ1050 Methanococcus jannaschii >gi 1499895 (U67548) conserved hypothetical protein [Methanococcus jannaschii]	2.5	
3490	AF035940	Homo sapiens MAGOH mRNA, complete cds	5e-096	3879018	(Z81108) similar to MAGO NASHI PROTEIN; cDNA EST yk415g7.3 comes from this gene; cDNA EST yk425g2.3 comes from this gene; cDNA EST yk425g2.5 comes from this gene; cDNA EST yk415g7.5 comes from this gene; cDNA EST yk415g7.5 comes from this gene; cDNA EST yk376g9.3 c	5e-027	
3491	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5e-013	3201662	(AF042191) paraxial protocadherin; PAPC [Danio rerio]	3.5	

		Nearest Neighbor			Nearest Neighbor			
200		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3492	S80107	membrane- associated diazepam binding inhibitor	e-113	244503	(S80107) membrane- associated diazepam binding inhibitor, MA-DBI [cattle, brain, Peptide, 552 aa] [Bos taurus]	2e-030		
3493	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4.00E-12	<none></none>	<none></none>	<none></none>		
3494	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-015	728834	!!!! ALU SUBFAMILY SB2 WARNING ENTRY	0.29		
3495	U32794	Haemophilus influenzae Rd section 109 of 163 of the complete genome	1.3	2369865	(Y14131) RNA polymerase [grapevine leafroll-associated virus 2]	5.1		
3496	AF030558	Rattus norvegicus phosphatidylinosi tol 5-phosphate 4- kinase gamma mRNA, complete cds	1e-013	<none></none>	<none></none>	<none></none>		
3497		Mouse mRNA for kinesin-like protein (Kif1b), complete cds	e-121	2497524	KINESIN-LIKE PROTEIN KIF1B mouse >gi 407339 gnl PI D d1005029 (D17577) Kif1b [Mus musculus]	1e-048		
3498		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-005		(Z73899) ZK829.5 [Caenorhabditis elegans]	1.5		
3499		Homo sapiens (subclone H8 5_a10 from P1 35 H5 C8) DNA sequence.	2e-018		(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.2		

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3500	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
3501	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
3502	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	
3503	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	2661842	(Y15732) DNA polymerase beta [Xenopus laevis]	4e-016	
3504	U65960	Human kinase substrate HASPP28 gene, 5' flanking region and partial cds	1e-021	2120084	reverse transcriptase - mouse >gi 558908	9.7	
3505	L19031	Rattus norvegicus organic anion transporter	3e-030		SODIUM- INDEPENDENT ORGANIC . ANION TRANSPORTER (ORGANIC ANION TRANSPORTING POLYPEPTIDE) anion - rat >gi 410311 (L19031) oatp [Rattus norvegicus]	2e-025	
3506		Homo sapiens beta-mannosidase mRNA, complete cds	0	3024091	BETA- MANNOSIDASE PRECURSOR beta-mannosidase [Homo sapiens]	2e-085	
3507		H.sapiens MICA gene	1e-055		hypothetical protein (L1H 3' region) - human	1e-009	

		Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)	•	(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3508	U50535	Human BRCA2 region, mRNA sequence CG006	4e-012	728831	IIII ALU SUBFAMILY J WARNING ENTRY	4.2		
3509	AF029984	Lycopersicon esculentum COP1 homolog (COP1) mRNA, complete cds	5e-035	3121867	COP1 REGULATORY PROTEIN sativum]	9e-052		
3510	Z59258	H.sapiens CpG DNA, clone 13d2, reverse read cpg13d2.rt1c	2e-046	3219914	HYPOTHETICAL 16.8 KD PROTEIN C30D10.04 IN CHROMOSOME II >gi 2276353 gn1 PI D e330328 pombe]	2e-009		
3511	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-008	<none></none>	<none></none>	<none></none>		
3512	AF004161	Oryctolagus cuniculus peroxisomal Ca- dependent solute carrier mRNA, complete cds	9e-030	2352427	(AF004161) peroxisomal Ca- dependent solute carrier	1e-025		
3513	U15643	Drosophila melanogaster ribosomal protein DL11 mRNA, complete cds	0.13	<none></none>	<none></none>	<none></none>		
3514	U15643	Drosophila melanogaster ribosomal protein DL11 mRNA, complete cds	0.13	<none></none>	<none></none>	<none></none>		
3515	X87212	H.sapiens mRNA for cathepsin C	e-103	1705632	DIPEPTIDYL- PEPTIDASE I PRECURSOR TRANSFERASE) >gi 2146949 pir  S 66504 dipeptidyl- peptidase I (EC 3.4.14.1) precursor - human sapiens]	3e-034		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3516	U28789	Mus musculus p53-associated cellular protein PACT mRNA, partial cds	e-101	<none></none>	<none></none>	<none></none>	
3517	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-009	127112	MAK16 PROTEIN >gi 73269 pir  BV BYK6 MAK16 protein - yeast (Saccharomyces cerevisiae) cerevisiae] >gi 595561 (U12980) Mak16p: putative nuclear protein [Saccharomyces cerevisiae]	5e-022	
3518	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	<none></none>	<none></none>	<none></none>	
3519	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	2650142	(AE001070) A. fulgidus predicted coding region AF0495	0.38	
3520	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	2500418	40S RIBOSOMAL PROTEIN S5 >gi 1203905	1.6	
3521	AF004161	Oryctolagus cuniculus peroxisomal Ca- dependent solute carrier mRNA, complete cds	9e-030	2352427	(AF004161) peroxisomal Cadependent solute carrier	1e-025	

		Nearest Neighbor			Nearest Neighbor			
250		lastN vs. Genbank)	T =	(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3522	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	121743	GTPASE-ACTIVATING PROTEIN (GAP) (RAS P21 PROTEIN ACTIVATOR) (P120GAP) (RASGAP) human >gi 182972 (M23379) GTPase-activating protein activating protein [Homo sapiens]	2.8		
3523	Z46372	R.norvegicus RNA for DNA topoisomerase II.	e-131	3876360	(Z68315) Similarity to Human MAP kinase phosphatase-1 (SW:PTN7_HUM AN) [Caenorhabditis elegans]	3e-011		
3524	X85060	B.taurus cosmid- derived microsatellite DNA	1e-051	2072972	(U93572) putative p150 [Homo sapiens]	1e-019		
3525	D86407	Homo sapiens DNA for apoER2, complete cds, and exon 19	0	3322933	(AE001238) DNA ligase (lig) [Treponema pallidum]	7.5		
3526	D17577	Mouse mRNA for kinesin-like protein (Kif1b), complete cds	e-130	2497524	KINESIN-LIKE PROTEIN KIF1B mouse >gi 407339 gnl PI D d1005029 (D17577) Kif1b [Mus musculus]	1e-049		
3527	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-007	2414623	(Z99259) putative phosphotransferas e	4e-009		
3528		Drosophila melanogaster strawberry notch (sno) mRNA, complete cds	1e-075	2076895	(AF002197) F20H11.2 gene product [Caenorhabditis elegans]	8e-057		

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3529	X54326	H.sapiens mRNA for glutaminyl- tRNA synthetase	0	135104	MULTIFUNCTIO NAL AMINOACYL- TRNA SYNTHETASE (CONTAINS: GLUTAMYL- TRNA SYNTHETASE glutamyl-prolyl- tRNA synthetase - human >gi 31958	3e-032	
3530	Z73360	Human DNA sequence from cosmid 92M18, BRCA2 gene region chromosome 13q12-13.	3e-039	2072955	(U93566) p40 [Homo sapiens]	7.8	
3531	Z73360	Human DNA sequence from cosmid 92M18, BRCA2 gene region chromosome 13q12-13.	3e-039	2072955	(U93566) p40 [Homo sapiens]	7.8	
3532	Z73360	Human DNA sequence from cosmid 92M18, BRCA2 gene region chromosome 13q12-13.	1e-040	2072955	(U93566) p40 [Homo sapiens]	0.012	
3533	Z73360	Human DNA sequence from cosmid 92M18, BRCA2 gene region chromosome 13q12-13.	1e-040	2072955	(U93566) p40 [Homo sapiens]	0.012	
3534	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	3808228	(AF039080) RNA dependent RNA polymerase [Sphaeropsis sapinea RNA virus 2]	1.5	

	ı	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3535	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
3536	U95760	Drosophila melanogaster strawberry notch (sno) mRNA, complete cds	3e-060	2078282	(U95760) Sno [Drosophila melanogaster]	1e-042	
3537	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.13	2832777	(AL021086) 1- evidence=predicte d by match; 1- match_accession= AA202870; 1- match_description =LD03215.5prime LD Drosophila melanogaster embryo BlueScript Drosophila melanogaster cDNA clone LD03215 5prime.; 1- match_species=Dr osop	4e-018	
3538	U95760	Drosophila melanogaster strawberry notch (sno) mRNA, complete cds	1e-075	2076895	(AF002197) F20H11.2 gene product [Caenorhabditis elegans]	8e-057	
3539	Z57610	H.sapiens CpG DNA, clone 187a10, reverse read cpg187a10.rt1a.	9e-061	913042	hepatocyte nuclear factor 3 beta, HNF3 beta	2e-014	
3540	X83416	H.sapiens PrP gene, exon 2	e-169	1172651	PROTEASE PRTH >gi 440338 (L27483) neutral protease large subunit [Porphyromonas gingivalis]	6.2	
3541	AF061016	Homo sapiens UDP-glucose dehydrogenase (UGDH) mRNA, complete cds	0	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.4	

		learest Neighbor		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	•	astN vs. Genbank)	DALLIE	ACCESSION DESCRIPTION P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE				
3542	X07290	Human HF.12 gene mRNA	7e-080	1127843	(U41164) Cys2/His2 zinc finger protein [Rattus norvegicus]	1e-034	
3543	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4.00E-12	<none></none>	<none></none>	<none></none>	
3544	U95760	Drosophila melanogaster strawberry notch (sno) mRNA, complete cds	3e-060	2078282	(U95760) Sno [Drosophila melanogaster]	1e-042	
3546	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.002	1255409	(U53153) similar to mouse bullous pemphigoid antigen, BPAG2 (PIR:A46053) [Caenorhabditis elegans]	7.3	
3547	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-008	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	9.9	
3548	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
3549	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	84605	glycine-rich protein GRP33 - brine shrimp	4.4	
3550	X83212	H.sapiens tryptophan hydroxylase gene, promoter region		807677	(M13101) unknown protein [Rattus norvegicus]	0.39	
3551	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.005	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
CEO	ACCESSION (B)	astN vs. Genbank)  DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
SEQ ID							
3552	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	310622	(L20249) homologous to Saccharopolyspora erythraea beta- ketoacyl synthase [Streptomyces coriofaciens]	0.4	
3553	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
3554	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-007	2996640	(AC004500) GDF- 9 [Homo sapiens]	8.2	
3555	Z35928	S.cerevisiae chromosome II reading frame ORF YBR059c	0.043	2384728	(AF015883) hydroxyproline- rich glycoprotein gas28p precursor [Chlamydomonas reinhardtii]	0.23	
3556	Z30174	M.domesticus (C57Bl/6J) mRNA for zinc finger protein 30	2e-037	543345	zinc finger protein 30 - mouse domesticus]	1e-020	
3557	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	<none></none>	<none></none>	<none></none>	
3558	M36072	Human ribosomal protein L7a (surf 3) large subunit mRNA, complete cds.	1e-054	133014	60S   RIBOSOMAL   PROTEIN L7A   (PLA-X   POLYPEPTIDE)   (SURF-3)   >gi 71116 pir  R5H   U7A ribosomal   protein L7a -   human   >gi 71117 pir  R5R   T7A ribosomal   protein L7a - rat   >gi 34203   (X52138) L7a   protein [Homo   sapiens] >gi 35512	0.019	

		Nearest Neighbor		Nearest Neighbor			
-		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					(X06705) PLA-X polypeptide [Homo sapiens]		
3559	U84720	Homo sapiens mRNA export protein (RAE1) mRNA, complete cds	2e-037	<none></none>	<none></none>	<none></none>	
3560	AE001054	Archaeoglobus fulgidus section 53 of 172 of the complete genome	1.2	<none></none>	<none></none>	<none></none>	
3561	U34683	Human glutathione synthetase mRNA, complete cds	3e-052	1346191	GLUTATHIONE SYNTHETASE (GLUTATHIONE SYNTHASE) (GSH SYNTHETASE) (GSH-S) sapiens] >gi 1236350 (U34683) glutathione synthetase	1e-014	
3562		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-015	1825695	(U88180) similar to molybdenum cofactor biosynthesis protein E [Caenorhabditis elegans]	4e-012	
3563		Plasmodium falciparum chromosome 2, section 58 of 73 of the complete sequence	0.005	<none></none>	<none></none>	<none></none>	

	<u> </u>	Nearest Neighbor		Nearest Neighbor			
1.4		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3564	D10871	Human h NAT allele 2-2 gene for arylamine N- acetyltransferase	5e-016	3915580	ZINC FINGER PROTEIN 186 finger protein [Homo sapiens]	0.96	
3565	M32251	Cat LINE-1 DNA sequence region 1.	2e-026	87765	hypothetical L1 protein (third intron of gene TS) - human >gi 364964 prf  15 10254A L1 repetitive element ORF [Homo sapiens]	2e-011	
3566	Y12773	H.sapiens TRIDENT/HFH1 1 gene, promoter sequence	3e-008	<none></none>	<none></none>	<none></none>	
3567	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	<none></none>	<none></none>	<none></none>	
3568	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	<none></none>	<none></none>	<none></none>	
3569	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-009	136821	HYPOTHETICAL PROTEIN UL13 precursor - human cytomegalovirus (strain AD169)	6	
3570		Homo sapiens caspase-activated nuclease mRNA, complete cds	e-104	3347857	(AF064019) DNA fragmentation factor 40 kDa subunit [Homo sapiens] >gi 3410909 gnl PI D d1033212 (AB013918) CAD	1e-024	
3571	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	2132458	probable membrane protein YDL211c - yeast	7.5	
3572	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	0.39	

	N	learest Neighbor		Nearest Neighbor				
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
ID								
		cds						
3573	U22233	Human	2e-015	2494053	5'-	0.02		
3313	022233	methylthioadenos			METHYLTHIOA			
		ine phosphorylase			DENOSINE			
		(MTAP) mRNA,		,	PHOSPHORYLA			
		complete cds.			SE (MTA			
					PHOSPHORYLA			
					SE) (MTAPASE) phosphorylase (EC			
					2.4.2.28) - human			
					>gi 847724			
				1	(U22233)			
					methylthioadenosi			
1					ne phosphorylase			
					[Homo sapiens]			
3574	X76122	A.majus cyclin-1	3.2	2135633	MHC cell surface	9		
		mRNA.			glycoprotein - human sapiens]			
	1105100	Y la avia	1e-009	699508	(U20542)	0.64		
3575	U95102	Xenopus laevis mitotic	16-009	099308	lethal(1)1Bi	0.0.		
		phosphoprotein			protein			
		90 mRNA,			[Drosophila			
		complete cds			melanogaster]	<u> </u>		
3576	D13391	Human CYP19	2e-018	<none></none>	<none></none>	<none></none>		
1		gene for			-			
1		aromatase			1			
		cytochrome P- 450, promoter		1				
		region						
		(containing two		1				
		cis-acting		İ				
		transcriptional						
1		regulatory						
		elements)	0.0000	522006	(III2975) C26E6 5	5e-045		
3577	U95102	Xenopus laevis mitotic	0.0002	532806	(U13875) C26E6.5 gene product	36-043		
		phosphoprotein			[Caenorhabditis			
		90 mRNA,			elegans]			
		complete cds						
3578	X63735	H.sapiens TRE5	4e-033	728831	!!!! ALU	9e-006		
		and TRE18			SUBFAMILY J			
		sequence of the			WARNING			
		tre oncogene			ENTRY	AIOVIE		
3579	AC004497	Homo sapiens	0.0005	<none></none>	<none></none>	<none></none>		
		chromosome 21,						
L	<u></u>	P1 clone				1		

		Nearest Neighbor			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	`	astN vs. Genbank)	DVALUE		DESCRIPTION	P VALUE		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	PVALUE		
ID_		LBNL#6						
3580	AB003095	Fruitfly strain SI259 mitochondrial DNA, A+T-rich region, partial sequence	0.12	<none></none>	<none></none>	<none></none>		
3581	Z36019	S.cerevisiae chromosome II reading frame ORF YBR150c	3.2	4107113	(AB007462) Pax- 2/5/8 [Ephydatia fluviatilis]	5.3		
3582	Z56421	H.sapiens CpG DNA, clone 117c7, reverse read cpg117c7.rt1a.	1e-033	3876101	(Z75536) similar to DnaJ domain; cDNA EST yk398h12.5 comes from this gene; cDNA EST yk250d6.5 comes from this gene [Caenorhabditis elegans]	1e-040		
3583	U36499	Human lymphoid-specific SP100 homolog (LYSP100-A) mRNA, complete cds	5e-015	1362890	phosphoprotein 75 - human >gi 402148	1e-008		
3584	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds		<none></none>	<none></none>	<none></none>		
3585	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	1045228	(X92429) Synthetase [Streptomyces anulatus]	0.84		
3586	D86963	Human mRNA for KIAA0208 gene, complete cds	0.04	<none></none>	<none></none>	<none></none>		
3587	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	<none></none>	<none></none>	<none></none>		

	1	Nearest Neighbor			Nearest Neighbor			
	` ` `	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3588	AB012113	Homo sapiens gene for CC chemokine PARC precursor, complete cds	0.0002	1723187	112.3 KD PROTEIN IN PYK1-SNC1 INTERGENIC REGION >gi 2131258 pir  S 70292 FUN12 protein Fun12p: 97kDa protein, function unknown [Saccharomyces cerevisiae]	4.2		
3589	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-007	<none></none>	<none></none>	<none></none>		
3590	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.002	<none></none>	<none></none>	<none></none>		
3591	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	6e-005	<none></none>	<none></none>	<none></none>		
3592	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.8		
3593	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-006	<none></none>	<none></none>	<none></none>		
3594	M80938	Oryza sativa 16.9 kDa heat shock protein gene, complete cds.	1.5	<none></none>	<none></none>	<none></none>		
3595	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-014	<none></none>	<none></none>	<none></none>		

	ľ	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3596	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	<none></none>	<none></none>	<none></none>	
3597	X67813	C.familiaris SRP72 mRNA for signal recognition particle	4e-083	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.2	
3598	AB007930	Homo sapiens mRNA for KIAA0461 perotein, partial cds	3e-038	3413884	(AB007930) KIAA0461 perotein [Homo sapiens]	3e-016	
3599	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	8e-007	3093586	(AF018165) amyloid precursor protein [Tetraodon fluviatilis]	2.7	
3600	Z35102	H.sapiens mRNA for Ndr protein kinase > :: emb A52140 A52 140 Sequence 6 from Patent WO9619579	e-126	2135799	Ndr protein kinase - human >gi 854170	9e-086	
3601	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
3602	X51544	Synthetic hamster-human hybrid cell (HCH-1) HSAG- 2 gene Alu repeat region.	0.13	1706266	SULFATE ADENYLATE TRANSFERASE SUBUNIT 2 (ATP- SULFURYLASE) >gi 1322409 gnl PI D e243270	5.8	
3603	Z98237	H.sapiens DNA for exon trapped sequence	3e-051	3979947	(AL034393) Y18D10A.15 [Caenorhabditis elegans]	6e-005	
3604	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	7e-005	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	,	lastN vs. Genbank)	_	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3605	M57465	N.crassa phytoene dehydrogenase (al-1) gene, complete cds.	0.29	<none></none>	<none></none>	<none></none>	
3606	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
3607	\$71335	Aox1=alternative oxidase {alternative pathway} suspension cells, mRNA, 1408 nt]	1.1	<none></none>	<none></none>	<none></none>	
3608	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.005	2621440	(AE000823) O- antigen transporter related protein	5.7	
3609	U36199	Caenorhabditis elegans CeMef-2 (mef-2) gene, complete cds.	1.1	259519	[(S48091) NSM [tomato spotted wilt virus TSWV, Peptide, 302 aa] [Tomato spotted wilt virus]	4.1	
3610	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.35	3399767	(U76298) uclacyanin I [Arabidopsis thaliana] >gi 3831466	0.35	
3611	AF000590	Homo sapiens chromosome 21q11-q21 genomic clone SA-292	7e-026	<none></none>	<none></none>	<none></none>	
3612	U64195	HIV-1 isolate ZP36 from Australia, reverse transcriptase (pol) gene, partial cds.	1.2	<none></none>	<none></none>	<none></none>	
3613	AB015331	Homo sapiens HRIHFB2017 mRNA, partial cds	1e-094	3970852	(AB015331) HRIHFB2017 [Homo sapiens]	0.0001	
3614	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

11	Nearest Neighbor			Nearest Neighbor			
	(B	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3615	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-009	1743885	(U79716) Human Reelin [Homo sapiens]	9.5	
3616	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-006	<none></none>	<none></none>	<none></none>	
3617	<none></none>	<none></none>	<none></none>	2338034	(AF005370) putative immediate early protein [Alcelaphine herpesvirus 1]	2e-008	
3618	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.006	2707286	(AF036316) cyclin [Prorocentrum minimum]	1.2	
3619	X79810	R.norvegicus CYP2C13 gene	0.049	2916892	(AL022004) PE_PGRS [Mycobacterium tuberculosis]	1	
3620	AJ224516	Gallus gallus IL-2 gene	1.4	<none></none>	<none></none>	<none></none>	
3621	Z79044	H.sapiens flow- sorted chromosome 6 HindIII fragment, SC6pA21C9	0.42	<none></none>	<none></none>	<none></none>	
3622	U39357	Ovis aries beta actin mRNA, complete cds	2e-024	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.3	
3623	U39357	Ovis aries beta actin mRNA, complete cds	1e-043	940346	(U20963) ORF1; late mRNA [Suid herpesvirus 1]	5.6	
3624	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	3e-008	2702361	(AF036706) No definition line found [Caenorhabditis elegans]	0.22	
3625	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.041	244874	Glvr-1 product [mice, Peptide, 681 aa]	1.9	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3626	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.002	. <none></none>	<none></none>	<none></none>	
3627	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.002	1730141	FRAGILE X MENTAL RETARDATION SYNDROME RELATED PROTEIN 2 >gi 2135129 pir  S 60173 fragile X mental retardation syndrome related protein - human >gi 1098637 (U31501) fragile X mental retardation syndrome related protein [Homo sapiens]	9.4	
3628	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-008	<none></none>	<none></none>	<none></none>	
3629	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	<none></none>	<none></none>	<none></none>	
3630	D87671	Rat mRNA for TIP120, complete cds	0	1799570	(D87671) TIP120 [Rattus norvegicus]	e-112	
3631	D87671	Rat mRNA for TIP120, complete cds	. 0	1799570	(D87671) TIP120 [Rattus norvegicus]	e-110	
3632	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3633	D88349	Chicken mRNA for tyrosinase, complete cds	0.12	2144081	luteinizing hormone/chorionic gonadotropin receptor - rat >gi 252167 bbs 10 9910 (S40803) luteinizing hormone/chorionic gonadotropin receptor, LH/CG receptor {alternatively spliced, clone rLHR1834}	9.3		
3634	X17206	Human mRNA for LLRep3	3e-025	2920827	(U92697) ribosomal protein S2 [Rattus norvegicus]	0.0003		
3635	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.3		
3636	X69878	H.sapiens Flt4 mRNA for transmembrane tyrosine kinase	2e-088	<none></none>	<none></none>	<none></none>		
3637	X69878	H.sapiens Flt4 mRNA for transmembrane tyrosine kinase	2e-088	<none></none>	<none></none>	<none></none>		
3638	X15509	Human gene for thymidine kinase, 5' region (EC 2.7.1.21)	4e-011	<none></none>	<none></none>	<none></none>		
3639	U89744	Rattus norvegicus putative cell surface antigen mRNA, complete cds	0.39	1085432	mucin (clone PGM-2A) - pig	0.0006		
3640	L29252	Human (clone D13-2) L-iditol-2 dehydrogenase gene, exon 4, exon 5, exon 6 and exon 7.	3e-006	83981	NADH dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 4 - Sauroleishmania tarentolae mitochondrion	2.4		

	N	Nearest Neighbor		Nearest Neighbor			
	(B)	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3641	Z35286	H.sapiens MDR3 gene, exon1, exon2	0.016	<none></none>	<none></none>	<none></none>	
3642	M11373	Simian T-cell leukemia virus, pol-env-pX-3' LTR region.	0.39	2773324	(AF040381) carbonic anhydrase [Erwinia carotovora]	5.9	
3643	M11373	Simian T-cell leukemia virus, pol-env-pX-3' LTR region.	0.39	2773324	(AF040381) carbonic anhydrase [Erwinia carotovora]	5.9	
3644	Z11763	O.granulifera gene for alpha- tubulin	0.39	2138321	(U89012) dentin matrix acidic phosphoprotein 1 [Homo sapiens]	2.6	
3645	<none></none>	<none></none>	<none></none>	1352944	HYPOTHETICAL 118.4 KD PROTEIN IN BAT2-DAL5 INTERGENIC REGION PRECURSOR YJR151c - yeast (Saccharomyces cerevisiae) >gi 1015903	3.9	
3646	U18351	Drosophila melanogaster insulin receptor gene, complete cds	0.005	1468983	(U64830) protein tyrosine kinase [Dictyostelium discoideum]	4e-012	
3647	M28458	Human growth hormone receptor gene, exon 2.	1.2	2648877	(AE000987) A. fulgidus predicted coding region AF1681	8.1	

	Nearest Neighbor			Nearest Neighbor			
	(B)	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3648	AF069139	HIV-1 isolate DH12 clone 5 from the USA, vpr protein (vpr) gene, partial cds; tat protein (tat) and rev protein (rev) genes, complete cds; vpu pseudogene, complete sequence; envelope glycoprotein (env) and nef protein (n	0.13	<none></none>	<none></none>	<none></none>	
3649	U42627	Rattus norvegicus tyrosine phosphatase mRNA, complete cds.	0.41	1070602	collagen alpha 1(II) chain precursor - human	0.55	
3650	Y12851	Homo sapiens P2X7 gene, exon 1 and joined CDS	0.005	<none></none>	<none></none>	<none></none>	
3651	U39706	Mycoplasma genitalium section 28 of 51 of the complete genome	0.39	465542	HYPOTHETICAL 20.0 KD PROTEIN IN TRNP 5'REGION (ORF160) >gi 625956 pir  S3 8599 hypothetical protein 160 (rpl20 5' region) - euglenid (Astasia longa) plastid	2	
3652	Z80361	H.sapiens HLA- DRB pseudogene, repeat region;	2e-048	<none></none>	<none></none>	<none></none>	
3653	U12171	Oryza sativa IR54 anther specific (RTS2) gene, complete cds.	3.5	<none></none>	<none></none>	<none></none>	
3654	AG001163	Homo sapiens genomic DNA, 21q region, clone: Q94A10X26	5e-014	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.004	
3655	X04780	Human tRNA- Tyr-pseudogene (clone pHtT2)	4.6	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3656	AF086264	Homo sapiens full length insert cDNA clone ZD43A10	0.002	<none></none>	<none></none>	<none></none>	
3657	AB011118	Homo sapiens mRNA for KIAA0546 protein, partial cds	0.002	1588661	tryptase [Bos taurus]	1.3	
3658	Z46379	Homo sapiens mRNA for anti- Sm antibody VH chain	0.13	<none></none>	<none></none>	<none></none>	
3659	Y12930	H.rustica CHD-W gene, intron	0.39	3861232	(AJ235272) PROBABLE TRANSPORT ATP-BINDING PROTEIN MSBA (msbA2) [Rickettsia prowazekii]	1.2	
3660	AF093267	Rattus norvegicus homer-1b mRNA, complete cds	0.005	<none></none>	<none></none>	<none></none>	
3661	M34057	Human transforming growth factor- beta 1 binding protein mRNA, complete cds.	0.043	<none></none>	<none></none>	<none></none>	
3662	X75418	H.sapiens TCR V Beta 13.2 gene (allele a).	0.4	<none></none>	<none></none>	<none></none>	
3663	Z68758	Human DNA sequence from cosmid cN85E10 on chromosome 22q11.2-qter	2e-025	3399771	(AF041839) Smad6 [Xenopus laevis]	0.39	
3664	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	2078282	(U95760) Sno [Drosophila melanogaster]	0.0006	
3665	Z75032	S.cerevisiae chromosome XV reading frame ORF YOR124c	0.14	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
OF O		astN vs. Genbank)	DIVALLE		s. Non-Redundant Pr DESCRIPTION	oteins) P VALUE	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION			
3666	U28831	Human protein immuno-reactive with anti-PTH polyclonal antibodies mRNA, partial cds. > :: gb I40055 I40055 Sequence 1 from patent US 5618695	0	896065	(U28831) protein that is immuno- reactive with anti- PTH polyclonal antibodies [Homo sapiens]	e-100	
3667	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.04	<none></none>	<none></none>	<none></none>	
3668	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
3669	Z96359	H.sapiens telomeric DNA sequence, clone 17QTEL013, read 17QTELOO013.s eq	7e-006	2921609	(AF039037) 980219 -this used to be part of R02C2.4 but was split into two genes based on protein similarities	7.7	
3670	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	8e-008	3342730	(AC005331) R31341_1 [Homo sapiens]	2e-019	
3671	U22460	Ictalurus punctatus heat shock protein 70 (CF Hsp70) mRNA, complete cds.	1.2	2143951	Ras-related protein - rat >gi 498257	5e-009	
3672	Y12259	R.norvegicus mRNA for Kir3.1 protein	0.005	135213	TYPE IIS RESTRICTION ENZYME ECO57I METHYLTRANS FERASE ACTIVITY >gi 281976 pir  S2 6426 type II site- specific deoxyribonuclease (EC 3.1.21.4) Eco57I endonuclease	9.9	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					[Escherichia coli]		
3673	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-008	3006154	(AL022299) putative cytochrome c1, heme protein precursor [Schizosaccharom yces pombe]	4.5	
3674	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	3915503	HYPOTHETICAL OXIDOREDUCT ASE IN CHEV- MOBA INTERGENIC REGION >gi 2632227 gnl PI D e1181911 1- dehydrogenase [Bacillus subtilis]		
3675	U71363	Human zinc finger protein zfp6 (ZF6) mRNA, partial cds	3e-070	2689441	(AC003682) F18547_1 [Homo sapiens]	4e-029	
3676	AF042275	Oryza sativa anther-specific protein gene, complete cds	0.39	<none></none>	<none></none>	<none></none>	
3677	M34601	P.berghei telomeric repeat region subfragment alpha DNA.	0.13	<none></none>	<none></none>	<none></none>	
3678	U09368	Human zinc finger protein ZNF140	6e-047	3445181	(AC005498) R31665_2 [Homo sapiens]	4e-027	
3679	D90345	Rat t complex polypeptide 1 (Tcp-1) mRNA	0.13	<none></none>	<none></none>	<none></none>	

	1	Nearest Neighbor		<u> </u>	Nagrast Najahhar	-
		lastN vs. Genbank)			Nearest Neighbor s. Non-Redundant P	roteine)
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
ID	15000550	1				
3680	AE000758	Aquifex aeolicus section 90 of 109 of the complete genome	0.38	134134	RYANODINE RECEPTOR, SKELETAL MUSCLE muscle - rabbit >gi 1710 (X15750) ryanodine receptor (AA 1-5037)	9.8
					[Oryctolagus cuniculus] >gi 1714 (X15209) ryanodine receptor [Oryctolagus cuniculus]	
3681	X60280	Vector plasmid pLTRpoly DNA	3e-040	2981631	(AB012223) ORF2 [Canis familiaris]	0.87
3682	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.8
3683	L81683	Homo sapiens (subclone 1_d11 from P1 H54) DNA sequence	3e-019	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	2
3684	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	9.7
3685	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.8
3686	X78261	H.sapiens mRNA for TRE17 5' extremity and unnamed adjacent to TRE17, locus tre-1.	3e-010	728836	!!!! ALU SUBFAMILY SP WARNING ENTRY	4.4
3687		Gallus gallus clone Ocyal unknown mRNA	1e-011		(AF093204) unknown [Gallus gallus]	0.097
3688		Homo sapiens (subclone H8 8_f5 from P1 35 H5 C8) DNA	3e-031		(U93570) p40 [Homo sapiens]	8e-006
			400	<u></u>		

	ı	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)			s. Non-Redundant P	roteins)	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		sequence.					
3689	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
3690	L10111	Octopus dofleini beta-tubulin mRNA, complete cds.	0.14	<none></none>	<none></none>	<none></none>	
3691	S83333	CYP27=sterol 27- hydroxylase/cere brotendinous xanthomatosis candidate gene {3' region, intron 6 to intron 8} [human, Genomic, 1725 nt, segment 4 of 4]	3.5	<none></none>	<none></none>	<none></none>	
3692	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.9	
3693	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	220578	(D00570) open reading frame (251 AA)	1.1	
3694	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	416563	INTESTINAL MEMBRANE A4 PROTEIN A4 differentiation- dependent protein [Homo sapiens]	0.021	
3695	AB018374	Mus musculus GARP34 mRNA, complete cds	4e-074	3724364	(AB018374) GARP34 [Mus musculus]	2e-017	
3696	AB018374	Mus musculus GARP34 mRNA, complete cds	4e-074	3724364	(AB018374) GARP34 [Mus musculus]	2e-017	
3697	AB013721	Oryctolagus cuniculus mRNA for mitsugumin 23, complete cds	4e-038	<none></none>	<none></none>	<none></none>	

	, in the second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
3698	U33147	Human mammaglobin mRNA, complete cds > :: gb I65735 I65735 Sequence 1 from patent US 5668267	1.1	1946371	(U93215) regulatory protein Viviparous-1 isolog	2.5		
3699	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0006	2132981	probable membrane protein YPL105c - yeast	5.1		
3700	U08802	HIV-1 sample 026 clone 06 from Thailand partial cds.	0.47	3880139	(Z68121) Similarity to Yeast nitrogen regulatory protein GLN3 (PIR Acc. No. S22280)	7.3		
3701	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-011	<none></none>	<none></none>	<none></none>		
3702	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	<none></none>	<none></none>	<none></none>		
3703	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	<none></none>	<none></none>	<none></none>		
3704	Z56740	H.sapiens CpG DNA, clone 13b5, reverse read cpg13b5.rt1c	4e-043	2465332	(U92819) unnamed HERV- H protein [Homo sapiens]	0.007		
3705	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-008	<none></none>	<none></none>	<none></none>		
3706	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	1293790	(U56248) Similar to polyketide synthase. [Caenorhabditis briggsae]	2.9		

	Ŋ	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3707	AF023283	Chikungunya virus S27 3'UTR	0.39	3560261	(AL031535) RNA binding protein	4.5	
3708	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
3709	AF030944	Brugia malayi microfilarial sheath protein SHP3a	0.12	<none></none>	<none></none>	<none></none>	
3710	AE000700	Aquifex aeolicus section 32 of 109 of the complete genome	0.15	<none></none>	<none></none>	<none></none>	
3711	AJ001050	Homo sapiens mRNA for thioredoxin reductase	4e-042	1843434	(D88687) KM- 102-derived reductase-like factor [Homo sapiens]	3e-038	
3712	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.002	625090	(U19464) outer arm dynein beta heavy chain [Paramecium tetraurelia] >gi 1588498 prf  2 208428A dynein:SUBUNIT =heavy chain [Paramecium tetraurelia]	2.7	
3713	AG001414	Homo sapiens genomic DNA, 21q region, clone: 9H11X4	0.46	<none></none>	<none></none>	<none></none>	
3714	AB007930	Homo sapiens mRNA for KIAA0461 perotein, partial cds	0	3413884	(AB007930) KIAA0461 perotein [Homo sapiens]	2e-068	
3715	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-007	<none></none>	<none></none>	<none></none>	
3716	Y09999	H.sapiens CHOP gene, intron 1	2e-007	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3717	AF023461	Homo sapiens FRA3B region sequence	0.13	2501500	ECDYSTEROID UDP- GLUCOSYLTRA NSFERASE PRECURSOR >gi 1563727 gnl PI D e267373 (Y08294) ecdysteroid UDP- glucosyltransferas e [Lacanobia oleracea granulovirus]	5.6	
3718	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	2330794	(Z98601) hypothetical protein	0.004	
3719	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	1363246	TIF1 protein - mouse >gi 998815 bbs 16 7126	5e-007	
3720	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	1314732	(U54640) 185 kDa silk protein [Chironomus pallidivittatus]	0.17	
3721	U09933	Human urokinase-type plasminogen receptor, exon 3	5e-025	3523099	(AF016271) Ksp- cadherin [Mus musculus]	7.6	
3722	M30187	S.cerevisiae mitochondrion Tyr-tRNA gene.	0.13	218437	(D90352) myo- inositol transporter	7.3	
3723	X79703	O.aries gene for beta-casein	0.043	141103	HYPOTHETICAL PROTEIN ORF- 1137 mouse	4.5	
3724		Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-009	2132008	hypothetical protein YOL072w - yeast	9.9	
3725	L39210	Homo sapiens inosine monophosphate dehydrogenase type II gene, complete cds	2e-078	2224711	(AB002383) KIAA0385 [Homo sapiens]	2e-018	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)	-	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3726	U52832	Homo sapiens Cri-du-chat region mRNA, clone CSC3	2e-005	<none></none>	<none></none>	<none></none>	
3727	AF015043	Homo sapiens EH-binding protein mRNA, partial cds	e-169	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.4	
3728	D28485	Human MSMB gene for beta- microseminoprote in (MSP), promoter region and exon1	4e-011	<none></none>	<none></none>	<none></none>	
3729	M33027	Human vasoactive intestinal peptide/PHM-27 gene, exons 1-6.	0.043	<none></none>	<none></none>	<none></none>	
3730	X15377	Human gene for the light and heavy chains of myeloperoxidase	2e-024	1346141	GLYCEROL KINASE (ATP:GLYCERO L 3- PHOSPHOTRAN SFERASE) (GLYCEROKINA SE) (GK) Mycoplasma genitalium (SGC3) >gi 3844648 (U39683) glycerol kinase (glpK) [Mycoplasma genitalium]	3e-011	
3731	X57103	Human h-lys gene for lysozyme (upstream region)	0.0005	<none></none>	<none></none>	<none></none>	
3732		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010		(AF077546) No definition line found [Caenorhabditis elegans]	9.8	
3733		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-012		(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3734	U83857	Human Aac11 (aac11) mRNA, complete cds	2e-027	2623755	(U35846) unknown [Mus musculus]	3e-005	
3735	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.5	
3736	U09367	Human zinc finger protein ZNF136	1e-065	1731412	ZINC FINGER PROTEIN 136 human >gi 487785 (U09367) zinc finger protein ZNF136	7e-060	
3737	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	2507475	PAIRED AMPHIPATHIC HELIX PROTEIN	5.8	
3738	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	3702452	(X80031) type IV collagen alpha 3 chain	1.5	
3739	AF086022	Homo sapiens full length insert cDNA clone YW23E02	3.5	<none></none>	<none></none>	<none></none>	
3740	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	2960225	(AL022120) PPE [Mycobacterium tuberculosis]	7.4	
3741	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
3742	AJ005866	Homo sapiens mRNA for putative Sqv-7- like protein, partial	e-177	4008517	(AJ005866) Sqv- 7-like protein [Homo sapiens]	9e-045	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3743	AF043231	Emericella nidulans cAMP- dependent protein kinase regulatory subunit (pkaR) gene, complete cds	1.1	<none></none>	<none></none>	<none></none>	
3744	AB002319	Human mRNA for KIAA0321 gene, partial cds	5e-066	2224583	(AB002319) KIAA0321 [Homo sapiens]	2e-024	
3745	M33132	Human proliferating cell nucleolar protein P120 gene, exons 1-15.	8e-018	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	0.077	
3746	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	2394463	(AF024498) No definition line found [Caenorhabditis elegans]	1.2	
3747	Z69944	S.pombe chromosome I cosmid c1F12.	4.4	<none></none>	<none></none>	<none></none>	
3748	Z81014	Human DNA sequence from cosmid U65A4, between markers DXS366 and DXS87 on chromosome X *	4e-022	896065	(U28831) protein that is immuno- reactive with anti- PTH polyclonal antibodies [Homo sapiens]	0.075	
3749	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	3877203	(Z70780) similar to initiation factor IF-2; cDNA EST CEMSD25F comes from this gene	4.4	
3750		Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>	
3751		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008		(Y12713) Pro-Pol- dUTPase polyprotein	2	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3752	M36072	Human ribosomal protein L7a (surf 3) large subunit mRNA, complete cds.	1e-054	133014	RIBOSOMAL PROTEIN L7A (PLA-X POLYPEPTIDE) (SURF-3) >gi 71116 pir  R5H U7A ribosomal protein L7a - human >gi 71117 pir  R5R T7A ribosomal protein L7a - rat >gi 34203 (X52138) L7a protein [Homo sapiens] >gi 35512 (X06705) PLA-X polypeptide [Homo sapiens]		
3753	AB001615	Homo sapiens DNA for cGMP- binding cGMP- specific phosphodiesteras e (PDE5), exon 1	6e-006	<none></none>	<none></none>	<none></none>	
3754	X57103	Human h-lys gene for lysozyme (upstream region)	5e-015	113670	!!!! ALU CLASS E WARNING ENTRY !!!!	3.3	
3755	L09708	Homo sapiens complement component 2 (C2) gene allele b, exons 10 through 18 and complete cds	6e-005	1143705	(X89760) Hox2a gene product [Zea mays]	9.7	
3756	X73685	C.aethiops hsp70 mRNA	2e-088	1322309	(U55176) heat shock cognate 70.II [Xenopus laevis]	2e-025	
3757	Z57594	H.sapiens CpG DNA, clone 186c5, reverse read cpg186c5.rt1b.	0.002	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3758	AF095927	Rattus norvegicus protein phosphatase 2C mRNA, complete cds	e-117	3777604	(AF095927) protein phosphatase 2C [Rattus norvegicus]	4e-040	
3759	U30788	Rattus norvegicus Tclone4 mRNA	5e-024	135576	LARGE TEGUMENT PROTEIN (VIRION PROTEIN UL36) >gi 73851 pir  WM BEH6 UL36 protein - human herpesvirus 1 (strain 17) >gi 59536 gnl PID  e312351 1]	1.6	
3760	Z96177	H.sapiens telomeric DNA sequence, clone 10QTEL040, read 10QTELOO040.s eq		1082626	myosin heavy chain VA - human (fragment)	5.8	
3761	M37463	E.gracilis chloroplast ribosomal protein genes rpl23, rpl2, rps19, rpl22, and rps3, complete cds.	0.38	2734883	(U75311) pyruvate decarboxylase 2 [Pichia stipitis]		
3762	AF086241	Homo sapiens full length insert cDNA clone ZD29F04	4e-064	3702137	(AL031393) dJ733D15.1 (Zinc- finger protein) [Homo sapiens]	1e-040	
3763	AF086241	Homo sapiens full length insert cDNA clone ZD29F04	4e-064	3702137	(AL031393) dJ733D15.1 (Zinc- finger protein) [Homo sapiens]	1e-040	
3764	AF008227	Drosophila melanogaster odd Oz product (odz) gene, exons 3, 4, 5, 6, 7, and complete cds	3.6	2661842	(Y15732) DNA polymerase beta [Xenopus laevis]	2e-020	
3765	AF039688	Homo sapiens antigen NY-CO-3 (NY-CO-3) mRNA, partial cds	0	3170176	(AF039688) antigen NY-CO-3 [Homo sapiens]	2e-073	

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3766	AF037332	Homo sapiens Eph-like receptor tyrosine kinase hEphB1b (EphB1) mRNA, complete cds	0.37	1255919	(X96511) MAFB protein [Coturnix japonica]	5.6	
3767	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	<none></none>	<none></none>	<none></none>	
3768	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.005	<none></none>	<none></none>	<none></none>	
3769	U95 <b>0</b> 94	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	<none></none>	<none></none>	<none></none>	
3770	X57103	Human h-lys gene for lysozyme (upstream region)	5e-015	113670	IIII ALU CLASS E WARNING ENTRY IIII	3.3	
3771	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	2947096	(U81032) TniQ [Pseudomonas stutzeri]	0.86	
3772	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	2947096	(U81032) TniQ [Pseudomonas stutzeri]	0.86	
3773	M84326	Human ADP- ribosylation factor 1 mRNA, complete cds	0	283748	collagen alpha 2(I) chain homolog - sea urchin (Strongylocentrotu s purpuratus) >gi 161436 purpuratus]	0.14	
3774	X82575	G.gallus mRNA for Cnot	0.39	3327136	(AB014561) KIAA0661 protein [Homo sapiens]	3e-033	

Nearest Neighbor Nearest Neighbor (BlastN vs. Genbank) (BlastX vs. Non-Redundant Proteins) ACCESSION SEO DESCRIPTION P VALUE ACCESSION DESCRIPTION P VALUE ID 3775 L43001 Bos taurus 3e-072 1730238 **GUANYLATE** 1e-030 guanylyl cyclase-**CYCLASE** activating protein **ACTIVATING PROTEIN 2** (GCAP 2) (RETINAL **GUANYLYL CYCLASE ACTIVATOR** PROTEIN P24) >gi|2136762|pir||A 57604 guanylate cyclase-activating protein 2 - bovine >gi|1002750 cyclase-activating protein 2 [Bos taurus] 3776 U47322 Cloning vector 7e-007 3335349 (AC004512) 9.2 DNA, complete Similar to sequence. gb|U46691 putative chromatin structure regulator (SUPT6H) from Homo sapiens. ESTs gb|T42908, gb|AA586170 and gb|AA395125 come from this gene. [Arabidopsis thaliana] 3777 L09647 Rattus norvegicus 2e-069 404764 (L10409) fork 3e-031 hepatocyte head related nuclear factor 3a protein [Mus musculus] 3778 U72756 Lycianthes 0.37 <NONE> <NONE> <NONE> heteroclita NADH dehydrogenase subunit protein, partial cds

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	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3779	M94314	Homo sapiens ribosomal protein L30 mRNA, complete cds	1e-073	3876073	(Z81505) similar to Zinc finger, C3HC4 type (RING finger); cDNA EST EMBL:D28025 comes from this gene; cDNA EST EMBL:D28024 comes from this gene; cDNA EST EMBL:D33210 comes from this gene; cDNA EST EMBL:D33241 comes from this	1.4	
3780	AF053315	Reporter vector pNFkB-Luc, complete sequence	9e-019	987050	(X65335) lacZ gene product [unidentified cloning vector]	0.3	
3781	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.5	
3782	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-013	1695957	(U78693) NADH dehydrogenase [Holmskioldia sanguinea]	1.9	
3783	AF074990	Homo sapiens full length insert cDNA YH85A11	0.005	1881709	(U89517) polyprotein [Dengue virus type 2]	9.6	
3784	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	<none></none>	<none></none>	<none></none>	
3785	AF020038	Homo sapiens NADP-dependent isocitrate dehydrogenase (IDH) mRNA, complete cds	4e-011	3647352	(Z97348) MAL3P1.11 [Plasmodium falciparum]	9.6	

	Nearest Neighbor			Nearest Neighbor			
	•	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3786	Z75199	S.cerevisiae chromosome XV reading frame ORF YOR291w	8e-028	3880560	(Z70271) Similarity to Yeast E1-E2 ATPase (SW:YED1_YEA ST); cDNA EST EMBL:D37634 comes from this gene; cDNA EST EMBL:D34653 comes from this gene; cDNA EST EMBL:D34527 comes from this gene; cDNA EST EMBL:D34527 comes from this	7e-048	
3787	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	<none></none>	<none></none>	<none></none>	
3788	M86400	Human phospholipase A2 mRNA, complete cds. > :: gb I34404 I34404 Sequence 8 from patent US 5597719	5e-088	<none></none>	<none></none>	<none></none>	
3789	X03100	Human HLA- SB(DP) alpha gene	0.47	3941737	(AF109719) BAT2 [Mus musculus]	2.4	
3790	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-015	3043662	(AB011141) KIAA0569 protein [Homo sapiens]	9.6	
3791	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	0.29	
3792	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	345555	Ig light chain - rainbow trout (fragment)	1.1	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3793	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
3794	AF064104	Homo sapiens Cdc14B1 phosphatase mRNA, complete cds	3e-030	2662463	(AF023158) tyrosine phosphatase [Homo sapiens]	1e-008	
3795	U29348	Salmonella enterica strain s2978 invasion protein SpaO (spaO), SpaP (spaP) and SpaQ (spaQ) genes, complete cds	0.0005	2291118	(AF016414) No definition line found [Caenorhabditis elegans]	9.6	
3796	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-016	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
3797	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	1168719	C6.1A PROTEIN	0.004	
3798	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	481236	hypothetical protein - Madagascar periwinkle roseus]	3.4	
3799	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-008	423157	finger protein ZNF33A - human (fragment)	4.3	
3800	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	
3801	U61950	Caenorhabditis elegans cosmid C45E5	1.2	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3802	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	1703028	CLATHRIN COAT ASSEMBLY PROTEIN AP47 HOMOLOG 2 (CLATHRIN COAT ASSOCIATED PROTEIN AP47 HOMOLOG 2) (GOLGI ADAPTOR AP-1 47 KD PROTEIN HOMOLOG 2) (HA1 47 KD SUBUNIT HOMOLOG 2) (CLATHRIN ASSEMBLY PROTEIN ASSEMBLY PROTEIN COMPL >gi 2134919 pir  A 57170 clathri	9.6	
3803	M31651	Homo sapiens sex hormone-binding globulin (SHBG) gene, complete cds	7e-017	<none></none>	<none></none>	<none></none>	
3804	D00596	Homo sapiens gene for thymidylate synthase, exons 1, 2, 3, 4, 5, 6, 7, complete cds	6e-038	<none></none>	<none></none>	<none></none>	
3805	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.7	
3806	D45906	Human mRNA for LIMK-2, complete cds	4e-096	<none></none>	<none></none>	<none></none>	
3807	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	2e-006	<none></none>	<none></none>	<none></none>	

1	Nearest Neighbor			Nearest Neighbor			
	(B)	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3808	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.004	<none></none>	<none></none>	<none></none>	
3809	AF045798	Xenopus laevis gremlin mRNA, complete cds	0.36	3551167	(AB012131) Ich1 [Coprinus cinereus]	4.1	
3810	D78275	Human mRNA for proteasome subunit p42, complete cds	8e-019	1709804	26S PROTEASE REGULATORY SUBUNIT S10B (P44) (CONSERVED ATPASE DOMAIN PROTEIN 44) 26S proteasome regulatory subunit [Homo sapiens]	0.001	
3811	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-009	<none></none>	<none></none>	<none></none>	
3812	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
3813	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-014	3193162	(AF067618) No definition line found [Caenorhabditis elegans]	1e-027	
3814	AF085858	Homo sapiens full length insert cDNA clone YN49B07	1e-017	3329465	(AF064553) NSD1 protein [Mus musculus]	4e-007	
3815	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-005	<none></none>	<none></none>	<none></none>	
3816	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0003	<none></none>	<none></none>	<none></none>	

	7	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3817	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-006	<none></none>	<none></none>	<none></none>	
3818	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2	<none></none>	<none></none>	<none></none>	
3819	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-006	416673	ATP SYNTHASE A CHAIN (PROTEIN 6) 3.6.1.34) protein 6 - liverwort (Marchantia polymorpha) mitochondrion >gi 786191 (M68929) atp6 [Marchantia polymorpha]	1.3	
3820	L14684	Rattus norvegicus nuclear-encoded mitochondrial elongation factor G mRNA, complete cds.	e-115	585084	ELONGATION FACTOR G, MITOCHONDRI AL PRECURSOR (MEF-G) >gi 543383 pir  S4 0780 translation elongation factor G, mitochondrial- rat >gi 310102	5e-038	
3821	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.2	
3822	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-012	1665789	(D87450) Similar to D.melanogaster parallel sister chromatids protein [Homo sapiens]	8.5	
3823	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-009	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
950		astN vs. Genbank)	D 1/4 1 1 1	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3824	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-015	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.2	
3825	L48489	Homo sapiens N- acetylglucosamin yltransferase III	1e-038	728831	IIII ALU SUBFAMILY J WARNING ENTRY	1e-008	
3826	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-008	<none></none>	<none></none>	<none></none>	
3827	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.9	
3828	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.4	
3829	AB012162	Homo sapiens mRNA for APC 2 protein, complete cds	1e-017	3894265	(AB012162) APC 2 protein [Homo sapiens]	0.45	
3830	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-010	1723680	HYPOTHETICAL 14.1 KD PROTEIN IN UPF3-SMD1 INTERGENIC REGION >gi 2132599 pir  S 64368 probable membrane protein YGR073c - yeast (Saccharomyces cerevisiae) >gi 1323101 gnl PI D e243468 (Z72858) ORF YGR073c [Saccharomyces cerevisiae]	1.3	

		learest Neighbor		1	Nearest Neighbor		
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID D							
3831	S54914	bup=5'of bmi-1	e-140	265569	(S54914) bup=5'of	2e-059	
		proviral insertion			bmi-1 proviral		
		locus [mice,			insertion locus [mice, Peptide,		
		Genomic/mRNA, 2022 nt]			195 aa] [Mus sp.]		
3832	U95102	Xenopus laevis	2e-009	<none></none>	<none></none>	<none></none>	
3632	093102	mitotic	20-007	TIONE	410112	110112	
		phosphoprotein					
		90 mRNA,					
		complete cds					
3833	U95094	Xenopus laevis	2e-012	<none></none>	<none></none>	<none></none>	
		XL-INCENP					
		(XL-INCENP)					
		mRNA, complete					
3834	U95102	cds Xenopus laevis	1e-015	2072296	(U95098) mitotic	2.9	
3634	093102	mitotic	10-015	2072250	phosphoprotein 44	_,,	
		phosphoprotein			[Xenopus laevis]		
		90 mRNA,					
		complete cds					
3835	U95094	Xenopus laevis	4e-007	1805270	(U81983)	6.6	
		XL-INCENP			endothelial PAS		
		(XL-INCENP) mRNA, complete			domain protein 1 [Mus musculus]		
		cds			[Mus musculus]		
3836	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
3837	X92814	H.sapiens mRNA	1e-032	1709969	H-REV 107	3e-013	
		for rat HREV107-			PROTEIN .		
		like protein			(1.5000000)	0.015	
3838	U95094	Xenopus laevis	1e-011	2183251	(AF002227) putative	0.015	
		XL-INCENP (XL-INCENP)			polyprotein		
		mRNA, complete			[border disease		
		cds			virus strain C413]		
3839	U95102	Xenopus laevis	1e-009	1142660	(U23502) POM1	7.3	
		mitotic			[Plasmodium		
1		phosphoprotein			chabaudi		
		90 mRNA,			chabaudi]		
2046	1105100	complete cds	4e-012	2072296	(U95098) mitotic	3.3	
3840	U95102	Xenopus laevis mitotic	46-012	2012290	phosphoprotein 44	د.د	
		phosphoprotein			[Xenopus laevis]		
		90 mRNA,			[han		
1		complete cds					
3841	U47322	Cloning vector	2e-058	224398	ORF [Simian virus	4e-005	
		DNA, complete			40]		
		sequence.	1				

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3842	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.9	
3843	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.3	
3844	Y15059	Homo sapiens hng/RC3 gene, exons 2,3 & 4	0.053	<none></none>	<none></none>	<none></none>	
3845	X99330	R.norvegicus mRNA for IP63 protein	2e-027	<none></none>	<none></none>	<none></none>	
3846	AF100303	Caenorhabditis elegans cosmid Y7G10A	0.53	<none></none>	<none></none>	<none></none>	
3847	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
3848	AF040094	Mus musculus inositol polyphosphate 5-phosphatase II (INPP5P) mRNA, complete cds	0.15	<none></none>	<none></none>	<none></none>	
3849	Y15724	Homo sapiens SERCA3 gene, exons 1-7 (and joined CDS)	2e-013	<none></none>	<none></none>	<none></none>	
3850	AB011144	Homo sapiens mRNA for KIAA0572 protein, partial cds	0	3043668	(AB011144) KIAA0572 protein [Homo sapiens]		
3851	AF020762	Homo sapiens clone 1400 unknown protein mRNA, partial cds	0	2738927	(AF020762) unknown protein [Homo sapiens]	2.8	
3852	Z99706	Human DNA sequence from cosmid U226D1 on chromosome X. Contains STS, complete sequence [Homo sapiens]	0.0002	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		`	s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3853	M73700	Human neutrophil lactoferrin mRNA, complete cds and 5' promoter region.	0.0002	<none></none>	<none></none>	<none></none>	
3854	D31793	Human CD40 ligand (CD40L) gene, 5' flanking region and exon 1	0.046	<none></none>	<none></none>	<none></none>	
3855	U16300	Human lysyl hydroxylase (PLOD) gene, intron 9, complete sequence.	0.0002	126363	LAMININ ALPHA-1 CHAIN PRECURSOR precursor - human	0.18	
3856	U61241	Homo sapiens p47-phox pseudogene, clone P41, exon 1	0.14	<none></none>	<none></none>	<none></none>	
3857	D37791	Mouse mRNA for beta-1,4- galactosyltransfer ase		3880102	(Z93390) similar to FYVE zinc finger; cDNA EST yk265b4.5 comes from this gene; cDNA EST yk359g9.5 comes from this gene; cDNA EST yk319c2.5 comes from this gene [Caenorhabditis elegans] zinc finger; cDNA EST yk265b4.5 comes from this gene; cDNA EST yk359g9	3e-021	
3858	Z57667	H.sapiens CpG DNA, clone 18a8, reverse read cpg18a8.rt1b.		<none></none>	<none></none>	<none></none>	
3859	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-014	2879809	(AJ223320) trp- like protein [Loligo forbesi]	1.5	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)			s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3860	U22296	Rattus norvegicus casein kinase 1 gamma 1 isoform mRNA, complete cds	e-126	3024053	CASEIN KINASE I, GAMMA 1 ISOFORM kinase 1 gamma 1 isoform [Rattus norvegicus]	1e-061	
3861	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-014	113669	III! ALU CLASS D WARNING ENTRY !!!!	2.6	
3862	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
3864	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	<none></none>	<none></none>	<none></none>	
3865	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	3182957	CGMP-INHIBITED 3',5'-CYCLIC PHOSPHODIEST ERASE B (CYCLIC GMP INHIBITED PHOSPHODIEST ERASE B) (CGI-PDE B) (CGIPDE1) phophodiesterase - human >gi 1145302 (U38178) cyclic nucleotide phophodiesterase [Homo sapiens] 3B [Homo sapiens]	4.4	
3866	AF099004	Caenorhabditis elegans cosmid C07D2	0.2	<none></none>	<none></none>	<none></none>	
3867	Z23091	H.sapiens GPV gene encoding platelet glycoprotein V precursor	5e-013	728836	!!!! ALU SUBFAMILY SP WARNING ENTRY	0.82	

		Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)			s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3868	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-007	2291255	(AF016430) weak similarity to Bacillus subtillis spore coat protein precursor (GB:L42066) and Dictyostelium discoideum calcium binding protein (NID:g426313) in proline-rich regions [Caenorhabditis elegans]	8.4	
3869	U58739	Caenorhabditis elegans cosmid F28C10.	0.33	<none></none>	<none></none>	<none></none>	
3870	L48473	Homo sapiens (subclone 7_e11 from P1 H16) DNA sequence.	3e-008	<none></none>	<none></none>	<none></none>	
3871	U95097	Xenopus laevis mitotic phosphoprotein 43 mRNA, partial cds	0.015	<none></none>	<none></none>	<none></none>	
3872	Z73360	Human DNA sequence from cosmid 92M18, BRCA2 gene region chromosome 13q12-13.	4e-020	<none></none>	<none></none>	<none></none>	
3873	Z71572	O.aries DNA for immunoglobulin joining regions	1.2	1699130	(U80027) weak similarity to Arabadopsis thaliana phytochrome E (PIR:S41912) [Caenorhabditis elegans]	6.1	
3874	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.7	

	7	Vearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3875	AB018263	Homo sapiens mRNA for KIAA0720 protein, partial cds	1.2	107240	oncogene 1 (tre-2 locus) (clone 210) - human	0.049	
3876	U87998	Mus musculus cyclin G1 gene, partial cds	0.14	<none></none>	<none></none>	<none></none>	
3877	AE001408	Plasmodium falciparum chromosome 2, section 45 of 73 of the complete sequence	1.8	<none></none>	<none></none>	<none></none>	
3878	AF061244	Agrocybe aegerita B type DNA polymerase (Mtpol) gene, complete cds; tRNA-Asn gene, complete sequence; and unknown genes, mitochondrial genes for mitochondrial products	0.16	3153241	(AF053004) class I cytokine receptor [Homo sapiens]	5.8	
3879	M73047	Homo sapiens tripeptidyl peptidase II mRNA, complete cds.	3e-028	136107	TRIPEPTIDYL- PEPTIDASE II (TPP II) tripeptidyl- peptidase II (EC 3.4.14.10) - human sapiens]	0.35	
3880	AB011393	Suncus murinus mitochondrial DNA, D-loop region, partial sequence, isolate TKU-M205	0.17	107422	proline-rich protein PRB3S (cys) - human	0.4	
3881	X69951	H.sapiens gene for casein kinase II alpha subunit > subunit alpha [human, Genomic, 18862 nt]	1e-008	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	0.54	

	ľ	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)			s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3882	U54558	Human translation initiation factor eIF3 p66 subunit mRNA, complete cds	3e-018	<none></none>	<none></none>	<none></none>	
3883	AB012259	Homo sapiens DNA, anonymous heat-stable fragment RP12-8	5e-012	<none></none>	<none></none>	<none></none>	
3884	U44130	Xenopus laevis p58 mRNA, partial cds	0.15	3873716	(Z74026) similar to 1- aminocyclopropan e-1-carboxylate synthase; cDNA EST EMBL:D34239 comes from this gene; cDNA EST EMBL:D35575 comes from this gene; cDNA EST EMBL:D64242 comes from this gene; cDNA EST EMBL:D67126 comes from 1- aminocyclopropan e-1-carbo	5.3	
3885	AB007917	Homo sapiens mRNA for KIAA0448 protein, complete cds	0.006	<none></none>	<none></none>	<none></none>	
3886	AJ223824	Lycopersicon esculentum cv Red River unknown sequence PCR random amplified RAPD band 9	0.045	<none></none>	<none></none>	<none></none>	
3887	U47322	Cloning vector DNA, complete sequence.	3e-008	2183251	(AF002227) putative polyprotein [border disease virus strain C413]	0.67	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID			4 006	AIONIES	<none></none>	<none></none>	
3888	U95102	Xenopus laevis mitotic	4e-006	<none></none>	\NONE>	NONE	
		phosphoprotein					
		90 mRNA,					
		complete cds				0.72	
3889	U67564	Methanococcus	1.3	2920535	(AF018081) type	0.73	
		jannaschii section			XVIII collagen [Homo sapiens]		
		106 of 150 of the complete genome			[Homo sapiens]		
3890	AE000720	Aquifex aeolicus	1.3	<none></none>	<none></none>	<none></none>	
3890	AE000720	section 52 of 109	1.5	110112			
		of the complete					
		genome					
3891	AB011230	Zaglossus bruijni	3.6	<none></none>	<none></none>	<none></none>	
		mitochondrial					
		gene for NADH					
		dehydrogenase subunit 1, partial					
		cds					
3892	Z96177	H.sapiens	1e-042	987050	(X65335) lacZ	0.0001	
		telomeric DNA			gene product		
ļ		sequence, clone			[unidentified		
		10QTEL040, read			cloning vector]		
		10QTELOO040.s					
3893	AF067646	eq Cloning vector	3e-029	987050	(X65335) lacZ	0.001	
3075	111 007010	pCMV-scriptEX,			gene product		
		complete			[unidentified		
		sequence			cloning vector]	AIONES	
3894	Z69919	Human DNA	3.8	<none></none>	<none></none>	<none></none>	
		sequence from cosmid 91K3,					
		Huntington's					
		Disease Region,					
		chromosome					
		4p16.3 contains					
		CpG island.		700110	COMITOTIC	10.012	
3895	X75757	G.gallus cycB3 mRNA.	6e-036	729112	G2/MITOTIC- SPECIFIC	4e-013	
		mKINA.			CYCLIN B3		
3896	L27833	Bos taurus	0.48	854348	(X87336) DNA	7.5	
		pregnancy-			endonuclease		
		associated			[Peperomia		
		glycoprotein-1			polybotrya]	1	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)			s. Non-Redundant Pr	oteins)	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3897	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.14	3169059	(AL023704) weak similarity to B.subtilis spore outgrowth factor B [Schizosaccharom yces pombe]	5e-052	
3898	X64123	H.sapiens PVR gene for poliovirus receptor (exon 8)	7e-006	2444416	(AF020484) NADH dehydrogenase- like protein [Gleditsia fera]	0.55	
3899	Z81043	Caenorhabditis elegans cosmid C29F3, complete sequence [Caenorhabditis elegans]	0.44	266459	P-SELECTIN PRECURSOR (GRANULE MEMBRANE PROTEIN 140) (GMP-140) (PADGEM) (CD62P) mouse >gi 200553 (M87861) P- selectin [Mus musculus]	1.8	
3900	AJ001235	Papio hamadryas ERV-9 like LTR insertion	3e-050	3126961	(AF061747) cell division protein FtsZ homolog	1.2	
3901	AE001314	Chlamydia trachomatis section 41 of 87 of the complete genome	1.2	<none></none>	<none></none>	<none></none>	
3902	X82895	H.sapiens mRNA for DLG2		3659505	(AC005084) similar to mouse mCASK-A; similar to e1288039	1e-054	
3903	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds		<none></none>	<none></none>	<none></none>	
3904	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	436923	(U01849) ORF1 [Trypanosoma brucei]	0.08	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
3905	D88982	Clostridium	0.38	1082769	RNA helicase A -	5.6	
		botulinum DNA			human		
		for C2 toxin					
		component-I and					
		component-II, complete cds					
3906	D50418	Mouse mRNA for	1e-041	2137398	homeotic protein	0.044	
3900	D30416	AREC3, partial	10-0-1	2137370	AREC3 (clone	0.044	
		cds			SM) - mouse		
3907	U95094	Xenopus laevis	6e-005	<none></none>	<none></none>	<none></none>	
		XL-INCENP					
		(XL-INCENP)					
		mRNA, complete					
	T105:01	cds		00115	(450006:2)	0.00	
3908	U95102	Xenopus laevis mitotic	2e-007	2314677	(AE000648) cation-transporting	0.36	
		phosphoprotein			ATPase, P-type		
		90 mRNA,			(copA)		
		complete cds			(CC). 2)		
3909	U72745	Dictyostelium	0.014	<none></none>	<none></none>	<none></none>	
		discoideum					
		cysteine					
2010	1 1011050	proteinase	2 221	NONE:	A LOND	3103 TE:	
3910	AJ011972	Homo sapiens mRNA for	3e-081	<none></none>	<none></none>	<none></none>	
		histone					
		deacetylase-like					
		protein (JM21)					
3911	U95094	Xenopus laevis	3e-011	<none></none>	<none></none>	<none></none>	
		XL-INCENP					
		(XL-INCENP)					
		mRNA, complete					
3912	U95102	Xenopus laevis	0.002	<none></none>	<none></none>	<none></none>	
1912	073102	mitotic	0.002	1101112	1101112	-110111	
		phosphoprotein					
		90 mRNA,					
		complete cds					
3913	AC001032	Homo sapiens	9e-009	130402	RETROVIRUS-	3.2	
		(subclone 2_c11			RELATED POL		
		from P1 H48) DNA sequence			POLYPROTEIN		
3914	J04830	S.cerevisiae	3.3	<none></none>	<none></none>	<none></none>	
3714	304030	CBP3 protein	ر.ر	71101112	-14OIAE	-1101112	
		gene, complete					
		cds.				ļ	
					L		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		`	s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3915	D78572	House mouse; Musculus domesticus mRNA for membrane glycoprotein, complete cds > :: dbj E12950 E129 50 cDNA GA3- 43 encoding novel polypeptide which appear when differentiate from embryo-tumor cell P19 to nerve cell	4e-044	1545807	(D78572) membrane glycoprotein [Mus musculus]	1e-020	
3916	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.1	
3917	U29923	Human AMP deaminase (AMPD3) gene, intron 1a and promoter 1b.	0.04	3256504	(AP000001) 115aa long hypothetical protein [Pyrococcus horikoshii]	0.094	
3918	Z68327	Human DNA sequence from cosmid U25D11, between markers DXS366 and DXS87 on chromosome X.	5e-015	<none></none>	<none></none>	<none></none>	
3919	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-013	<none></none>	<none></none>	<none></none>	
3920	M89955	Human 5-HT1D- type serotonin receptor gene, complete cds.	0	112819	5- HYDROXYTRYP TAMINE 1D RECEPTOR human > gi 177772 (M89955) 5- HT1D-type serotonin receptor receptor:ISOTYPE =1D-alpha [Homo	3e-053	

	<u> </u>	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID					sapiens]		
3921	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete	2e-008	3879698	(Z78065) predicted using Genefinder	9.1	
3922	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.002	3184285	(AC004136) hypothetical protein [Arabidopsis thaliana]	9.5	
3923	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.005	139805	XFIN PROTEIN >gi 65234 (X06021) Xfin protein (AA 1 - 1350) [Xenopus laevis]	1.9	
3924	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	<none></none>	<none></none>	<none></none>	
3925	AF013711	Homo sapiens 22 kDa actin-binding protein	1e-020	103509	I factor 2 (transposon) - fruit fly protein [Drosophila teissieri]	5.5	
3926	S83526	red photopigment gene {Alu repeat region, long intron 1} [human, peripheral blood leucocytes, Genomic, 1987 nt]	7e-006	<none></none>	<none></none>	<none></none>	
3927	AB011542	Homo sapiens mRNA for MEGF9, partial cds	0	3449310	(AB011542) MEGF9 [Homo sapiens]	2e-095	

		learest Neighbor		Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3928	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
3929	X67312	P.pijperi mitochondrion DNA for Vaccinia virus- like terminal loop structure	6e-006	<none></none>	<none></none>	<none></none>	
3930	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	3080474	(AL022602) cell divisin protein FtsW	1.2	
3931	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-006	3769486	(AF074946) DNA polymerase [hemorrhagic enteritis virus]	1.3	
3932	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	<none></none>	<none></none>	<none></none>	
3933	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	1890266	(U88585) NADH- dehydrogenase subunit 1 [Quedius mesomelinus]	4.2	
3934	Z12112	pWE15A cosmid vector DNA	1e-051	987050	(X65335) lacZ gene product [unidentified cloning vector]	4e-009	
3935	AF023180	Listeria monocytogenes low temperature requirement A protein (ltrA) gene, complete cds	0.005	<none></none>	<none></none>	<none></none>	
3936	D10856	D. melanogaster cyclin A gene	0.37	2315521	(AF016452) similar to the beta transducin family	1e-028	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)	:	(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
3937	U95102	Xenopus laevis	9e-009	3687507	(AL031788) C2H2	7.3	
		mitotic			type zinc finger		
		phosphoprotein			protein [Schizosaccharom		
		90 mRNA,			yces pombe]		
2000	700061	complete cds	2e-078	<none></none>	<none></none>	<none></none>	
3938	Z80361	H.sapiens HLA- DRB pseudogene,	2e-078	NONE/	\I\O\I\L>	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	
		repeat region;					
3939	L22551	Plasmodium	1.2	<none></none>	<none></none>	<none></none>	
3939	L22331	yoelii yoelii	1.2	NONE	TONE	1.01.2	
		merozoite surface					
		protein 1 gene, 5'					
		end.					
3940	X74178	B.taurus	0.005	2291118	(AF016414) No	2.5	
		microsatellite			definition line		
		DNA INRA153			found		
					[Caenorhabditis		
					elegans]	0.40	
3941	U95094	Xenopus laevis	9e-010	1354361	(U52008) Mrp50	0.48	
		XL-INCENP			[Streptococcus		
	!	(XL-INCENP)		•	pyogenes]		
		mRNA, complete					
3942	U41635	Human OS-9	0.12	<none></none>	<none></none>	<none></none>	
3342	041033	precurosor	0.12	110112			
		mRNA, complete					
		cds					
3943	U95102	Xenopus laevis	1e-011	<none></none>	<none></none>	<none></none>	
		mitotic					
		phosphoprotein					
		90 mRNA,					
		complete cds				4.2	
3944	M37470	Human beta-N-	5e-025	728832	!!!! ALU SUBFAMILY SB	4.3	
		acetylhexosamini dase (HEXB)			WARNING		
		gene, deletion			ENTRY		
		junction.			LIVIKI		
3945	U95094	Xenopus laevis	6e-006	97885	salivary agglutinin	0.84	
	0,50,74	XL-INCENP			receptor precursor		
		(XL-INCENP)			- Streptococcus		
		mRNA, complete			sanguis		
		cds					

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3946	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	140550	HYPOTHETICAL 259 KD PROTEIN (ORF 2136) >gi 81341 pir  A05 037 hypothetical protein 2136 - liverwort (Marchantia polymorpha) chloroplast >gi 11665	2.5	
3947	L13176	Papio anubis apolipoprotein C- I gene, partail mRNA.	0.0005	<none></none>	<none></none>	<none></none>	
3948	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-006	580702	(X74410) fixP gene product [Azorhizobium caulinodans]	2.9	
3949	X92987	B.primigenius mRNA for coat protein gamma- cop	2e-036	1706000	COATOMER GAMMA SUBUNIT (GAMMA-COAT PROTEIN) (GAMMA-COP) >gi 1066165 (X92987) coat protein gamma- cop [Bos primigenius]	2e-008	
3950	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-008	223232	protein src [Avian sarcoma virus]	0.37	
3951	AF037350	Rattus norvegicus NF-E2-related factor 2 mRNA, complete cds	1e-013	3004573	(AC004520) similar to NFE2- related transcription factors; similar to I48694 (PID:g2137676) [Homo sapiens]	8e-073	
3952	AJ011972	Homo sapiens mRNA for histone deacetylase-like protein (JM21)	8e-092	<none></none>	<none></none>	<none></none>	

	N	learest Neighbor		Nearest Neighbor			
	<b>,</b> ,	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID				NO. IE	A IONES	AIONES	
3953	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
3954	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
3955	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	630444	CR5 protein - Trypanosoma brucei >gi 468424	4.3	
3956	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	630444	CR5 protein - Trypanosoma brucei >gi 468424	4.3	
3957	AF086172	Homo sapiens full length insert cDNA clone ZB89E10	9e-062	1172991	60S RIBOSOMAL PROTEIN L21 sapiens] >gi 984143 (X89401) ribosomal protein L21 [Homo sapiens] >gi 1096939 prf  2 113200B ribosomal protein L21	9e-024	
3958	D42084	Human mRNA for KIAA0094 gene, partial cds	2e-058	1703270	PUTATIVE METHIONINE AMINOPEPTIDA SE 1 (METAP 1) (PEPTIDASE M 1) (KIAA0094) product is related to S.cerevisiae methionine aminopeptidase. [Homo sapiens]	1e-016	

	ľ	Nearest Neighbor		Nearest Neighbor			
	(B)	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3959	AF034755	Homo sapiens microphthalmia- associated transcription factor (MITF) gene, promoter region and partial cds	2e-005	<none></none>	<none></none>	<none></none>	
3960	Z96177	H.sapiens telomeric DNA sequence, clone 10QTEL040, read 10QTELOO040.s eq		<none></none>	<none></none>	<none></none>	
3961	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	141028	NADH- UBIQUINONE OXIDOREDUCT ASE CHAIN 5 >gi 76351 pir  QQ UTC5 NADH dehydrogenase (ubiquinone)	1.1	
3962	U93237	Human menin (MEN1) gene, complete cds	0.37	134853	TRANSCRIPTIO N INITIATION PROTEIN SPT5 yeast (Saccharomyces cerevisiae) >gi 172680 (M62882) SPT5 protein [Saccharomyces cerevisiae] >gi 854480 (Z49810) Spt5p [Saccharomyces cerevisiae]	0.49	
3963	Z93782	Caenorhabditis elegans cosmid R12G8, complete sequence [Caenorhabditis elegans]	0.008	1171084	A/G-SPECIFIC ADENINE GLYCOSYLASE	6.5	
3964	U11270	Human antithrombin III gene, exon 1 and partial cds.	2e-023	728837	!!!! ALU SUBFAMILY SQ WARNING ENTRY	9e-006	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3965	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-007	3650488	(AF042273) signal transducing adaptor molecule 2A [Homo sapiens]	3.6	
3966	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
3967	AF086207	Homo sapiens full length insert cDNA clone ZC48C05	1e-009	1077301	probable membrane protein YOL101c - yeast similarity with bee NADH- ubiquinone oxidoreductase chain 2 [Saccharomyces cerevisiae] >gi 1419955 gnl PI D e252291	0.41	
3968	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.2	
3969	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	2274853	(AJ000502) iron regulatory protein	0.15	
3970	U82165	Cercopithecus aethiops transmembrane glycoprotein CD99-cos7 mRNA, partial cds	2e-015	2735010	(U82166) CD99 type II-COS7 [Cercopithecus aethiops]	0.011	
3971	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3972	M87680	Human simple repeat polymorphism.	3e-040	3874946	(Z79598) cDNA EST EMBL:D34748 comes from this gene; cDNA EST yk218e6.5 comes from this gene; cDNA EST yk244e3.5 comes from this gene; cDNA EST yk248a4.5 comes from this gene; cDNA EST yk250a3.5 comes from this gene; cDNA EST	1e-008	
3973	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	119396	ENV POLYPROTEIN (COAT POLYPROTEIN) reticuloendothelios is virus >gi 61786 (X01455) env- protein (capsid protein) [Reticuloendotheli osis virus] >gi 209712 (K02537) envelope polyprotein [Avian reticuloendothelios is virus A]	4.6	
3974	AB011143	Homo sapiens mRNA for KIAA0571 protein, complete cds	e-151	1708199	HSC70- INTERACTING PROTEIN	4e-023	
3975	AC001050	Homo sapiens (subclone 3_e9 from P1 H55) DNA sequence	1e-019	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	3e-006	
3976	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-008	1077543	probable membrane protein YDR198c - yeast	5.9	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3977	AJ005175	Drosophila virilis mRNA for GAGA factor class B-isoform	0.056	<none></none>	<none></none>	<none></none>	
3978	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-014	478731	replication protein - Butyrivibrio fibrisolvens plasmid pRJF1 >gi 152515 (M94552) replication protein [Plasmid pRJF1]	1.5	
3979	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-006	3319480	(AF077546) No definition line found [Caenorhabditis elegans]	6.5	
3980	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3	
3981	AF003350	Mus musculus Npc1 gene, and npc-nih intron containing the MaLR inserted sequence	4e-007	1170261	OUTER MEMBRANE USHER PROTEIN HIFC PRECURSOR	6.4	
3982	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.001	<none></none>	<none></none>	<none></none>	
3983	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-006	<none></none>	<none></none>	<none></none>	
3984	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-007	<none></none>	<none></none>	<none></none>	
3985	AB007939	Homo sapiens mRNA for KIAA0470 protein, complete cds	e-163	3413902	(AB007939) KIAA0470 protein [Homo sapiens]	2e-057	

	ì	Nearest Neighbor		Nearest Neighbor			
	•	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3986	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.9	
3987	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-006	<none></none>	<none></none>	<none></none>	
3988	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1e-006	<none></none>	<none></none>	<none></none>	
3989	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.008	2414527	(Z99263) hypothetical protein MLCB637.01c [Mycobacterium leprae]	1.3	
3990	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.074	464237	NADH- UBIQUINONE OXIDOREDUCT ASE CHAIN 4	2.2	
3991	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010		(Z69360) Weak similarity to . Eimeria thrombospondin (PIR Acc. No. A45517); cDNA EST EMBL:M89266 comes from this gene; cDNA EST yk295b9.5 comes from this gene [Caenorhabditis elegans] Eimeria thrombospondin (PIR Acc. No. A45517); cDNA EST EMBL:M89266 comes	7.7	

		Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
3992	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	400624	SODIUM- AND CHLORIDE- DEPENDENT GABA TRANSPORTER 2 >gi 348413 pir  A4 5078 gamma- aminobutyric acid transporter protein 2 - rat >gi 202523 (M95762) GABA transporter [Rattus norvegicus]	0.62	
3993	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-015	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.9	
3994	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.9	
3995	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	2286159	(AF007831) glycoprotein H [Human herpesvirus 7]	6.3	
3996	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-014	<none></none>	<none></none>	<none></none>	
3997	D16888	Human HepG2 3' region cDNA, clone hmd2c03	e-104	<none></none>	<none></none>	<none></none>	
3998	U00995	Rattus norvegicus TA1 mRNA, complete cds.	1e-031	3639058	(AF077866) amino acid transporter E16 [Homo sapiens]	1e-050	

	N	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
ID						~ ~ ~		
3999	AF037219	Homo sapiens PIX1 mRNA sequence	5e-013	586863	HYPOTHETICAL 9.2 KD PROTEIN IN RECR-BOFA INTERGENIC REGION >gi 1075824 pir  A 41869 bofA 5'- region hypothetical protein orf74 - Bacillus subtilis subtilis] >gi 2632289 gnl PI D e1181955	2.7		
4000	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-009	<none></none>	(Z99104) yaaL <none></none>	<none></none>		
4001	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-015	<none></none>	<none></none>	<none></none>		
4002	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-013	549734	HYPOTHETICAL 105.7 KD PROTEIN IN TPK3-PIR1 INTERGENIC REGION >gi 481105 pir  S3 7786 hypothetical protein YKL165c - yeast (Saccharomyces cerevisiae) >gi 407483 (Z26877) unknown [Saccharomyces cerevisiae] >gi 486289 (Z28165) ORF YKL165c	3e-019		

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4003	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA,	2e-014	<none></none>	<none></none>	<none></none>	
4004	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-008	228110	T cell receptor variable region:SUBUNIT =beta:ISOTYPE=1 9 [Rattus norvegicus]	3.6	
4005	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-014	930045	(X15332) alpha-1 (III) collagen [Homo sapiens]	0.52	
4006	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.2	
4007	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-015	2960195	(Y13051) tax [Human T-cell lymphotropic virus type 2b]		
4008	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-007	3523099	(AF016271) Ksp- cadherin [Mus musculus]	6.6	
4009	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-015	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.7	
4010	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-015	<none></none>	<none></none>	<none></none>	
4011	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	2121280	(AF000270) lipoprotein [Borrelia burgdorferi]	1.5	

	N	learest Neighbor		Nearest Neighbor			
	,	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4012	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	<none></none>	<none></none>	<none></none>	
4013	L20489	Zea mays NADH dehydrogenase subunit 4 (complex I) (nad4) gene, exon 4.	3.5	<none></none>	<none></none>	<none></none>	
4014	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-014	<none></none>	<none></none>	<none></none>	
4015	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-016	927407	(X89858) actin binding protein [Drosophila melanogaster]	0.02	
4016	U05659	Human 17beta- hydroxysteroid dehydrogenase type 3 mRNA, complete cds	1e-092	1169300	ESTRADIOL 17 BETA- DEHYDROGENA SE 3 DEHYDROGENA SE) >gi 1085271 pir  S 43928 17-beta- hydroxysteroid dehydrogenase - human >gi 531162 hydroxysteroid dehydrogenase:IS OTYPE=3 [Homo sapiens]	4e-029	
4017	U02428	Cloning vector pDR2, complete sequence	2e-066	987050	(X65335) lacZ gene product [unidentified cloning vector]	2e-009	

		Nearest Neighbor			Nearest Neighbor			
	<b>`</b>	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4018	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-018	3979938	(AL034393) predicted using Genefinder; cDNA EST yk343c12.5 comes from this gene; cDNA EST yk402e12.5 comes from this gene; cDNA EST yk457e8.5 comes from this gene; cDNA EST yk470f2.5 comes from this gene; cDNA EST yk470f2.5 comes	7e-009		
4019	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	<none></none>	<none></none>	<none></none>		
4020	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	804806	(M13100) unknown protein [Rattus norvegicus]	5.7		
4021	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-014	<none></none>	<none></none>	<none></none>		
4022	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	<none></none>	<none></none>	<none></none>		
4023	U49974	Human mariner2 transposable element, complete consensus sequence	e-124	1698455	(U49974) mariner transposase [Homo sapiens]	2e-028		
4024	L31840	Rattus norvegicus nuclear pore complex protein NUP107 mRNA, complete cds.	e-175	1709212	NUCLEAR PORE COMPLEX PROTEIN NUP107	3e-093		

	N	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4025	AB001632	Homo sapiens DNA for cGMP- binding cGMP- specific phosphodiesteras e (PDE5), exon	7e-007	<none></none>	<none></none>	<none></none>	
4026	X96401	H.sapiens mRNA for ROX protein	8e-070	<none></none>	<none></none>	<none></none>	
4027	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.015	<none></none>	<none></none>	<none></none>	
4028	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.2	
4029	AJ006064	Rattus norvegicus mRNA for coronin-like protein	e-124	3757680	(AJ006064) coronin-like protein [Rattus norvegicus]	2e-091	
4030	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	1184072	(U40766) COL-1 [Meloidogyne incognita]	0.019	
4031	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.002	231721	T-CELL SURFACE GLYCOPROTEIN CD8 ALPHA CHAIN PRECURSOR (T- LYMPHOCYTE DIFFERENTIATI ON ANTIGEN T8/LEU-2) >gi 38145 (X60223) CD8 alpha chain	5.8	
4032	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-009	<none></none>	<none></none>	<none></none>	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4033	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.005	<none></none>	<none></none>	<none></none>	
4034	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-011	1020391	(L48340) alcohol dehydrogenase [Methylobacteriu m extorquens]	1.4	
4035	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	2291282	(AF016433) similar to C. elegans olfactory receptor ODR-10 (NID:g1235900) [Caenorhabditis elegans]	4.4	
4036	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	478993	DNA-binding protein TAF-II 250K - fruit fly TATA-binding protein associated factor II 250, TBP associated factor II 250, TAFII250 {C-terminal}	5e-006	
4037	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.4	
4038	X03100	Human HLA- SB(DP) alpha gene	2e-025	<none></none>	<none></none>	<none></none>	
4039	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-008	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID	102700	Human	2e-048	3874988	(Z74029)	5.6	
4040	J03798	autoantigen small	26-046	3074900	Similarity to	3.0	
		nuclear			C.elegans alcohol		
		ribonucleoprotein			dehydrogenase		
		Sm-D mRNA,			(WP:C17G10.8);		
		complete cds.			cDNA EST	1	
		1			EMBL:D66106		
					comes from this		
					gene; cDNA EST		
					EMBL:D69117		
	:				comes from this		
				ļ	gene; cDNA EST EMBL:D69761		
					comes from this		
					gene; cDNA EST		
					EMBL:C12156		
					come		
4041	U95102	Xenopus laevis	6e-006	2292986	(AJ000496) cyclic	0.5	
		mitotic		`	nucleotide-gated		
1		phosphoprotein			channel beta		
		90 mRNA,			subunit [Rattus		
		complete cds	2 010	2072206	norvegicus]	2.5	
4042	U95102	Xenopus laevis	3e-010	2072296	(U95098) mitotic phosphoprotein 44	2.3	
		mitotic phosphoprotein			[Xenopus laevis]		
		90 mRNA,			[Menopus lastis]		
		complete cds					
4043	AF020187	Amblyomma	1.2	<none></none>	<none></none>	<none></none>	
		americanum			-		
		ecdsyteroid					
		receptor			NONE	AVONTES	
4044	Z68758	Human DNA	2e-035	<none></none>	<none></none>	<none></none>	
		sequence from cosmid cN85E10					
		on chromosome					
		22q11.2-qter					
4045	U95102	Xenopus laevis	8e-008	2529632	(L78917) virion	4.6	
		mitotic			protein [Rubella		
1		phosphoprotein			virus]		
		90 mRNA,					
		complete cds	1	AIO) TO	AIONES	AIONES	
4046	U95102	Xenopus laevis	6e-005	<none></none>	<none></none>	<none></none>	
		mitotic					
1	1	phosphoprotein 90 mRNA,					
		complete cds					
L		complete cus	<u> </u>	<u> </u>	<u> </u>		

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID			- 016	700001	I I I I I I I I I I I I I I I I I I I	0.063	
4047	AB007957	Homo sapiens mRNA, chromosome 1 specific transcript KIAA0488	2e-016	728831	!!!! ALU SUBFAMILY J WARNING ENTRY		
4048	M64716	Human ribosomal protein S25 mRNA, complete cds.	3e-082	2660720	(AF029678) PHF1 [Homo sapiens]	7e-013	
4049	AB002437	Homo sapiens mRNA from chromosome 5q21-22, clone:LI33	6e-026	<none></none>	<none></none>	<none></none>	
4050	Z74 <b>8</b> 93	S.cerevisiae chromosome XV reading frame ORF YOL151w	0.13	<none></none>	<none></none>	<none></none>	
4051	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-006	<none></none>	<none></none>	<none></none>	
4052	U43416	Human replication control protein 1 (PARC1) mRNA, complete cds.	2e-056	728831	IIII ALU SUBFAMILY J WARNING ENTRY	0.007	
4053	AF042346	Homo sapiens putative phenylalanyl- tRNA synthetase beta-subunit mRNA, complete cds	0	4104933	(AF042346) putative phenylalanyl- tRNA synthetase beta-subunit; PheHB [Homo sapiens]	e-123	
4054		<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4055	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4056		Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
4057	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0005	2981221	(AF053091) eyelid [Drosophila melanogaster]	2.6	

	Ŋ	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4058	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
4059	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-007	<none></none>	<none></none>	<none></none>	
4060	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	9.6	
4061	U11081	Human type 1 vasoactive intestinal peptide receptor (V1RG) gene, exon 3.	0.43	<none></none>	<none></none>	<none></none>	
4062	X82272	Human endogenous retrovirus env mRNA	8e-081	1196429	(M14123) pol/env ORF (bases 3878- 8257) first start codon at 4172; Xxx; putative [Homo sapiens]	6e-058	
4063	S61789	NF1=neurofibro matosis type 1 {deletion breakpoint, tetrameric STR} [human, neurofibrosarcom a tissue, Genomic Mutant, 698 nt]	0.0005	2494294	NEUROGENIC LOCUS NOTCH 3 PROTEIN	4.3	
4064	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0005	3264773	(AF072439) zinc- finger protein-37; ZFP-37 [Rattus norvegicus]	3.3	
4065	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.5	
4066	U47322	Cloning vector DNA, complete sequence.	9e-054	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	0.6	

	<u> </u>	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4067	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.4	
4068	U31557	Ovine adenovirus IVa2 protein gene, DNA polymerase gene, terminal protein gene and 52,55 kDa protein gene, partial cds	0.0002	3002875	(AF042104) envelope glycoprotein [Human immunodeficiency virus type 1]	2.6	
4069	AL023973	Human DNA sequence from clone 1033E15 on chromosome 22q13.1-13.2. Contains part of a novel gene, ESTs and a GSS, complete sequence [Homo sapiens]	7e-017	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.061	
4070	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	2e-005	<none></none>	<none></none>	<none></none>	
4071	X07679	Xenopus laevis XK70A gene for type I keratin	0.39	2281044	(Z95636) laminin alpha 5 chain [Homo sapiens]	0.9	
4072	X96886	H.sapiens spcDNA, clone 2- 65	5e-014	<none></none>	<none></none>	<none></none>	
4073	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	9e-008	<none></none>	<none></none>	<none></none>	
4074	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	6e-005	1079278	activin receptor II STK3 precursor - African clawed frog >gi 260044 bbs 11 8656 (S49438) activin receptor, XAR1 [Xenopus, oocytes, Peptide,	1.3	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID					510 aa]		
					310 aaj		
4075	AF097909	Peptostreptococc	0.046	<none></none>	<none></none>	<none></none>	
		us micros fibril-					
		like structure					
		subunit FibA					
		(fibA) gene,					
		complete cds; excreted protein					
		FibB (fibB) gene,					
		partial cds; and					
		unknown gene					
4076	U95094	Xenopus laevis	4e-010	<none></none>	<none></none>	<none></none>	
		XL-INCENP					
		(XL-INCENP)					
		mRNA, complete		1			
1077	AL009008	cds Plasmodium	0.45	<none></none>	<none></none>	<none></none>	
4077	ALUU9UU8	falciparum DNA	0.43	\INOINE>	NONE	TIOINE	
		***					
•		SEQUENCING					
		IN PROGRESS					
		*** from contig					
1		3-58, complete					
1050	7.04606	sequence	0.015	AIONES	NONE	<none></none>	
4078	L34686	Serpulina byodysantoriae	0.015	<none></none>	<none></none>	NOINE	
		hyodysenteriae flagellar protein					
4079	AJ130718	Homo sapiens	1e-022	3582136	(AB015432)	2e-008	
70/)	13150710	mRNA for			LAT1 (L-type		
		glycoprotein-			amino acid		
		associated amino			transporter 1)		
		acid transporter			[Rattus		
		y+LAT1		NGS	norvegicus]	AIO) III:	
4080	X51969	Cyprinus carpio	1.2	<none></none>	<none></none>	<none></none>	
		growth hormone					
4001	1105004	gene Xenopus laevis	3e-010	2072296	(U95098) mitotic	1.2	
4081	U95094	XL-INCENP	76-010	2012290	phosphoprotein 44	1.4	
		(XL-INCENP)			[Xenopus laevis]		
		mRNA, complete		I			
		cds					
	<u> </u>	cds		<u> </u>	<u> </u>		

		learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID	11002001011						
4082	U95094	Xenopus laevis	4e-011	2072296	(U95098) mitotic	3.4	
		XL-INCENP			phosphoprotein 44		
		(XL-INCENP)			[Xenopus laevis]		
		mRNA, complete					
		cds					
4083	L38961	Human putative	1e-071	1174470	OLIGOSACCHA	1e-008	
		transmembrane			RYL		
		protein precursor			TRANSFERASE		
		(B5) mRNA,			STT3 SUBUNIT		
		complete cds			HOMOLOG (B5)		
					(INTEGRAL MEMBRANE		
					PROTEIN 1)		
					musculus]		
					>gi 1588285 prf  2		
					208301A integral		
			1		membrane protein		
					[Mus musculus]		
4084	U95102	Xenopus laevis	2e-013	267449	HYPOTHETICAL	7e-014	
1004	0,5102	mitotic			12.5 KD		
		phosphoprotein			PROTEIN		
		90 mRNA,			ZK637.2 IN		
		complete cds			CHROMOSOME		
		_			III		
					>gi 102507 pir  S1		
					5787 hypothetical		
					protein 1 (cosmid		
					ZK637) -		
					Caenorhabditis		
					elegans Genefinder; cDNA		
					EST yk217b5.3		
					comes from this		
					gene; cDNA EST		
					yk217b5.5 comes		
				1	from this gene;		
1					cDNA EST		
				1	yk340g12.3		
4085	U95098	Xenopus laevis	8e-008	2072296	(U95098) mitotic	3.4	
		mitotic			phosphoprotein 44		
		phosphoprotein			[Xenopus laevis]		
		44 mRNA, partial					
		cds			3,63,55	210272	
4086	X77733	T.aestivum	0.005	<none></none>	<none></none>	<none></none>	
		VDAC 1 mRNA.		<u> </u>	<u> </u>		

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4087	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	3123172	ZINC FINGER PROTEIN 151 (MIZ-1 PROTEIN) >gi 2230871 gnl PI D e286602 (Y09723) Miz-1 protein [Homo sapiens]	2e-010	
4088	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-010	180498	(M17517) complement H factor [Homo sapiens]	5.8	
4089	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-007	<none></none>	<none></none>	<none></none>	
4090	U24697	Chironomus samoensis nanos homolog (Cs nos) gene, complete cds.	0.13	3880999	(AL021492) Y45F10D.11 [Caenorhabditis elegans]	7e-022	
4091	Ü95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.7	
4092	U81504	Homo sapiens beta-3A-adaptin subunit of the AP-3 complex mRNA, complete cds	6e-088	2199512	(U81504) beta- 3A-adaptin subunit of the AP- 3 complex [Homo sapiens]	0.0001	
4093	AF053304	Homo sapiens mitotic checkpoint component Bub3	e-108	3378104	(AF047473) testis mitotic checkpoint BUB3 [Homo sapiens]	3e-024	
4094	S70431	type-1 angiotensin II receptor {exons 1 and 2, promoter} [human, peripheral lymphocytes, Genomic, 2853 nt, segment 1 of 2]	4e-013	126295	LINE-1 REVERSE TRANSCRIPTAS E HOMOLOG	3e-005	

	N	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)	:	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4095	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-013	<none></none>	<none></none>	<none></none>	
4096	D10355	Human mRNA for alanine aminotransferase	3e-082	111345	alanine transaminase (EC 2.6.1.2) - rat	4e-042	
4097	AF043252	Homo sapiens mitochondrial outer membrane protein (Tom40) gene, nuclear gene encoding mitochondrial protein, exons 7, 8 and 9	e-167	3941342	(AF043250) mitochondrial outer membrane protein [Homo sapiens] >gi 3941347 (AF043253) mitochondrial outer membrane protein [Homo sapiens] >gi 4105703 (AF050154) D19S1177E [Homo sapiens]	7e-013	
4098	U41668	Human deoxyguanosine kinase mRNA, complete cds	e-125	2833282	DEOXYGUANOS INE KINASE PRECURSOR sapiens]	2e-009	
4099	AF017416	Bacillus thuringiensis d- endotoxin gene, complete cds	0.14	<none></none>	<none></none>	<none></none>	
4100	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-008	<none></none>	<none></none>	<none></none>	
4101	AF017416	Bacillus thuringiensis d- endotoxin gene, complete cds	0.14	<none></none>	<none></none>	<none></none>	
4102	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-008	<none></none>	<none></none>	<none></none>	
4103	AJ003081	Homo sapiens repetitive DNA	5e-024	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4104	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-007	<none></none>	<none></none>	<none></none>	
4105	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-006	1572756	(U70848) C43G2.1 gene product [Caenorhabditis elegans]	4e-038	
4106	U33915	Craterostigma plantagineum myb-related transcription factor (cpm10) gene, complete cds	0.14	<none></none>	<none></none>	<none></none>	
4107	U46493	Cloning vector pFlp recombinase gene, complete cds	5e-033	987050	(X65335) lacZ gene product [unidentified cloning vector]	0.004	
4108	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-009	3417298	(AC002044) Alpha-fetoprotein enhancer binding protein (3' partial) [Homo sapiens]	0.33	
4109	M16039	Dictyostelium discoideum pst- cath gene encoding pst- cathepsin, complete cds.	0.0002	<none></none>	<none></none>	<none></none>	
4110	D21851	Human mRNA for KIAA0028 gene, partial cds	6e-005	<none></none>	<none></none>	<none></none>	
4111	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-007	1723920	HYPOTHETICAL 37.4 KD PROTEIN IN SEC27-SSM1B INTERGENIC REGION >gi 2131603 pir  S 64149 hypothetical protein YGL136c - yeast (Saccharomyces cerevisiae) >gi 1246842 gnl PI	8e-006	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID					D e210737 (X92670) G2830		
4112	X75861	H.sapiens TEGT gene	e-180	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.6	
4113	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-008	1399962	(U62317) choline kinase isolog 384D8_3 [Homo sapiens]	0.67	
4114	Y07660	M.tuberculosis accBC gene	2e-059	465847	HYPOTHETICAL 66.5 KD PROTEIN F02A9.5 IN CHROMOSOME III >gi 280542 pir  S2 8313 hypothetical protein F02A9.5 - Caenorhabditis elegans Genefinder; similar to Propionyl-CoA carboxylase beta chain; cDNA EST EMBL:M89018 comes from this gene; cDNA EST EMBL:D2806	4e-056	
4115	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.014	765086	(D30786) feline CD9 [Felis catus]	1.9	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
4116	D29011	Human mRNA	e-125	2136006	proteasome	4e-008	
		for proteasome			subunit MB1 -		
		subunit X,			human (fragment) MB1=LMP7		
		complete cds			homolog [human,		
					JY T-cells, Peptide		
					Partial, 215 aa]		
					[Homo sapiens]		
4117	U95094	Xenopus laevis	4e-012	2072296	(U95098) mitotic	6.2	
		XL-INCENP			phosphoprotein 44		
		(XL-INCENP)			[Xenopus laevis]		
		mRNA, complete					
	711600	cds	e-178	119172	ELONGATION	6e-054	
4118	Z11692	H.sapiens mRNA for elongation	e-1/8	119172	FACTOR 2 (EF-2)	06-034	
		factor 2			eEF-2 - human		
		factor 2			>gi 31106		
					(X51466)		
					elongation factor 2		
		-			factor 2 [Homo		
					sapiens]		
4119	AF070530	Homo sapiens	0	3387886	(AF070530)	4e-013	
		clone 24751			unknown [Homo sapiens]		
4120	D12646	unknown mRNA Mouse kif4	6e-057	1170659	KINESIN-LIKE	2e-022	
4120	D12040	mRNA for	00-037	1170037	PROTEIN KIF4	20 022	
		microtubule-			musculus]		
		based motor			-		
		protein KIF4,	i		•		
		complete cds			110117	3.103.175	
4121	U95094	Xenopus laevis	1e-009	<none></none>	<none></none>	<none></none>	
		XL-INCENP					
		(XL-INCENP) mRNA, complete					
		cds					
4122	X75861	H.sapiens TEGT	e-180	2072296	(U95098) mitotic	2.6	
		gene			phosphoprotein 44		
L					[Xenopus laevis]		
4123	U95094	Xenopus laevis	4e-012	630864	LRR47 protein -	0.0002	
		XL-INCENP			fruit fly		
		(XL-INCENP)			(Drosophila melanogaster)		
		mRNA, complete					
		cus			(X75760) LRR47		
					[Drosophila		
					melanogaster]		

		Nearest Neighbor		Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	` `	astN vs. Genbank)		` `			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4124	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5e-014	3876775	(Z81077) predicted using Genefinder; Similarity to Yeast protein 8248 (TR:G587531)	1e-015	
4125	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-008	480516	transposase (clone 22.5) - African malaria mosquito transposon mariner (fragment) >gi 159600	2.8	
4126	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-008	<none></none>	<none></none>	<none></none>	
4127	X65279	pWE15 cosmid vector DNA	2e-066	987050	(X65335) lacZ gene product [unidentified cloning vector]	4e-015	
4128	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.8	
4129	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-008	<none></none>	<none></none>	<none></none>	
4130	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-008	<none></none>	<none></none>	<none></none>	
4131	X74871	H.sapiens gene for RNA pol II largest subunit, exons 20-22	1.1	1182038	(Z69368) unknown [Schizosaccharom yces pombe]	0.86	
4132	M64983	Human fibrinogen beta chain gene, complete mRNA. > gb I47706 I47706 Sequence 3 from patent US 5639940	0.23	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID		1:64	6 057	1170650	KINESIN-LIKE	2e-022	
4133	D12646	Mouse kif4	6e-057	1170659	PROTEIN KIF4	26-022	
		mRNA for microtubule-			musculus]		
		based motor			musculusj		
		protein KIF4,					
		complete cds					
4134	D86957	Human mRNA	1.1	<none></none>	<none></none>	<none></none>	
	200,0,	for KIAA0202					
		gene, partial cds					
4135	U95094	Xenopus laevis	7e-006	<none></none>	<none></none>	<none></none>	
		XL-INCENP					
		(XL-INCENP)					
		mRNA, complete					
1100	1 (2000)	cds	4e-008	<none></none>	<none></none>	<none></none>	
4136	M20902	Human apolipoprotein C-	46-006	NONE>	NONE	-IVOIVE	
		I (VLDL) gene,					
		complete cds.					
4137	L36849	Cloning vector	9e-040	987050	(X65335) lacZ	9e-007	
		pZEO (isolate			gene product		
		SV1)			[unidentified		
		phleomycin/zeoci			cloning vector]		
		n-binding protein					
		gene, complete					
4120	V90010	cds.	0	<none></none>	<none></none>	<none></none>	
4138	X80910	H.sapiens PPP1CB mRNA				0.23	
4139	M77812	Rabbit myosin	0.0002	2088793	(AF003150) - similar to cuticular	0.23	
		heavy chain mRNA, complete			collagen		
		cds.			[Caenorhabditis		
	İ	Cus.			elegans]		
4140	U41165	Human	0.13	<none></none>	<none></none>	<none></none>	
1		recombination					
		'hot spot' region					
		associated with					
		the CMT1A					
		duplication and the HNPP					
		deletion					
1		containing a	1				
		mariner					
		transposon-like					
		element			3763	210315	
4141	U95094	Xenopus laevis	0.0006	<none></none>	<none></none>	<none></none>	
		XL-INCENP					
		(XL-INCENP) mRNA, complete		ľ			
		cds					
L		1	1		I	J	

	N	learest Neighbor		1	Nearest Neighbor	
		astN vs. Genbank)		(BlastX vs	s. Non-Redundant Pro	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
4142	AC001502	Homo sapiens (subclone 2_c7 from P1 H43) DNA sequence	0.014	3164130	(D78600) cytochrome P450 monooxygenase	7.5
4143	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>
4144	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
4145	L31760	Human STS UT8178.	0.17	<none></none>	<none></none>	<none></none>
4146	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	<none></none>	<none></none>	<none></none>
4147	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0006	2662587	(AF036696) contains similarity to Brassica oleracea non-green plastid phosphate/triose- phosphate translocator precursor (GB:U13632) [Caenorhabditis elegans]	2e-016
4148	X56807	Human DSC2 mRNA for desmocollins type 2a and 2b	6e-037	319943	desmocollin 3b precursor - human	7e-014
4149	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-012	<none></none>	<none></none>	<none></none>
4150	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	2854155	(AF045640) contains similarity to ion channel proteins	3.4

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
4151	U95102	Xenopus laevis	3e-010	2507153	VACUOLAR	0.011	
		mitotic			PROTEIN		
		phosphoprotein			SORTING-		
		90 mRNA,			ASSOCIATED		
	,	complete cds			PROTEIN VPS16		
					>gi 2133204 pir  S		
					62031 vacuolar		
					protein sorting-		
					associated protein		
					VPS16 - yeast	ļ <b>l</b>	
1					(Saccharomyces		
ŀ					cerevisiae)		
			1		>gi 1171414	1	
					(U44030) Vsp16p:		
1					Vacuolar sorting		
		1			protein		
					[Saccharomyces cerevisiae]		
		1:04	2.025	3877579		2e-054	
4152	D12646	Mouse kif4	2e-035	38/13/9	(Z82271) Similarity to	26-034	
		mRNA for			Mouse kinensin-		
		microtubule-			like protein KIF4		
		based motor	Ì		(SW:P33174);		
		protein KIF4, complete cds			cDNA EST		
		complete cus			EMBL:D27320		
					comes from this		
-			1		gene; cDNA EST		
					EMBL:D27322		
					comes from this		
					gene; cDNA EST		
					EMBL:D27321		
					comes from this		
					gene; cDNA EST		
					EMBL:D35764		
1					comes Mouse		
					kinensin-like		
					protein		

	J	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
ID					(700071)	2 054		
4153	D12646	Mouse kif4	2e-035	3877579	(Z82271)	2e-054		
		mRNA for			Similarity to			
		microtubule-			Mouse kinensin-			
		based motor			like protein KIF4			
		protein KIF4,			(SW:P33174); cDNA EST	ŀ		
		complete cds			EMBL:D27320			
					comes from this			
					gene; cDNA EST			
					EMBL:D27322			
					comes from this			
					gene; cDNA EST			
					EMBL:D27321			
					comes from this			
			!		gene; cDNA EST			
					EMBL:D35764			
					comes Mouse			
					kinensin-like			
					protein			
4154	D12646	Mouse kif4	2e-035	3877579	(Z82271)	9e-058		
		mRNA for			Similarity to			
		microtubule-			Mouse kinensin-			
		based motor			like protein KIF4	j ,		
		protein KIF4,			(SW:P33174);			
		complete cds			cDNA EST EMBL:D27320	] ]		
1					comes from this			
					gene; cDNA EST			
					EMBL:D27322			
					comes from this	•		
					gene; cDNA EST			
					EMBL:D27321			
					comes from this			
					gene; cDNA EST			
1					EMBL:D35764			
					comes kinensin-			
					like protein KIF4			
4155	M30539	Human SK2 c-	0.13	137334	66 KD PROTEIN	10		
		Ha-ras-1			>gi 77357 pir  JQ0			
		oncogene-			107 hypothetical			
		encoded protein			66K protein -			
		gene, exon 1.			Ononis yellow mosaic virus			
L			1		mosaic virus	ll		

Nearest Neighbor Nearest Neighbor (BlastX vs. Non-Redundant Proteins) (BlastN vs. Genbank) DESCRIPTION P VALUE SEQ ACCESSION DESCRIPTION P VALUE ACCESSION ID 1173044 60S 3e-007 2e-086 4156 L05096 Homo sapiens **RIBOSOMAL** ribosomal protein **PROTEIN L39** L39 mRNA, norvegicus] complete cds >gi|1373419 (U57846) ribosomal protein L39 ribosomal protein L39 [Homo sapiens] 987050 (X65335) lacZ 0.18 Plasmid pKA1 2e-025 4157 D13749 gene product **DNA** [unidentified cloning vector] (Z95890) 6.3 2e-057 2131036 4158 AF007157 Homo sapiens PE PGRS clone 23856 [Mycobacterium] unknown mRNA. tuberculosis] partial cds 0.0008 (AB014577) AF031400 Poecilia orri 1.2 3327168 4159 KIAA0677 protein NADH [Homo sapiens] dehydrogenase subunit 2 gene, mitochondrial gene encoding mitochondrial protein, complete cds 3e-009 <NONE> <NONE> <NONE> 4160 U58468 Human vasoactive intestinal peptide gene, 5' flanking sequence from -5172 to -1924 1.2 4e-032 2119507 alpha-1C-D11078 Homo sapiens 4161 RGH2 gene, adrenergic receptor isoform 2 retrovirus-like - human element >gi|927209|gnl|PI D|d1007476 (D32202) alpha 1C adrenergic receptor isoform 2 [Homo sapiens] <NONE> <NONE> <NONE> <NONE> <NONE> <NONE> 4162 !!!! ALU 0.002 2e-023 728831 M31061 Human ornithine 4163 **SUBFAMILY J** decarboxylase WARNING gene, complete cds. **ENTRY** 

and the first of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of

		learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		`	s. Non-Redundant Pr DESCRIPTION	oteins) P VALUE	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION			
4164	M19980	M.fervidus gap gene encoding glyceraldehyde-3- phosphate dehydrogenase, complete cds.	0.4	1825606	(U88169) similar to molybdoterin biosynthesis MOEB proteins [Caenorhabditis elegans]	3e-057	
4165	D17036	Human HepG2 partial cDNA, clone hmd3e08m5	5e-025	<none></none>	<none></none>	<none></none>	
4166	L14714	C. elegans cosmid ZC97.	0.39	3874412	(Z70034) similarity to 35.1KD hypothetical yeast protein (Swiss Prot accession number P38805); cDNA EST CEMSE65F comes from this gene; cDNA EST EMBL:T01315 comes from this gene; cDNA EST yk452e10.3 comes from this gene; cDNA 35.1KD hypothetical yeast p		
4167	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-008	<none></none>	<none></none>	<none></none>	
4168		Caenorhabditis elegans cosmid C33D3, complete sequence [Caenorhabditis elegans]	0.044	3876784	(Z81530) predicted using Genefinder	5.9	
4169	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-010	3549676	(AL031394) putative protein	3.1	
4170	D87001	Human (lambda) DNA for immunoglobulin light chain	0.36	3133246	(AB013170) NADH dehydrogenase subunit 5	2.4	

		Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)  ACCESSION   DESCRIPTION   P VALUE				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION				
4171	M37191	Human ras inhibitor mRNA, partial cds.	e-122	107561	Ras inhibitor (clone JC310) - human sapiens]	3e-035		
4172	AB018374	Mus musculus GARP34 mRNA, complete cds	2e-046	3724364	(AB018374) GARP34 [Mus musculus]	2e-008		
4173	X62527	R.norvegicus gene for CNS- myelin proteolipid protein (exon 6)	1.2	1155068	(X94976) cell wall-plasma membrane linker protein	1.6		
4174	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-008	2781355	(AC003113) F24O1.11 [Arabidopsis thaliana]	0.52		
4175	AF002715	Homo sapiens MAP kinase kinase kinase (MTK1) mRNA, complete cds	e-168	2352277	(AF002715) MAP kinase kinase kinase [Homo sapiens]	1e-042		
4176	U07807	Human metallothionein IV (MTIV) gene, complete cds.	0.047	<none></none>	<none></none>	<none></none>		
4177	D11129	Pneumonia virus of mice gene 7	0.14	<none></none>	<none></none>	<none></none>		
4178	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>		
4179	AF070557	Homo sapiens clone 24422 mRNA sequence	0	<none></none>	<none></none>	<none></none>		
4180	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.005	<none></none>	<none></none>	<none></none>		
4181	AF045765	Homo sapiens G protein-coupled receptor	9e-018	728833	IIII ALU SUBFAMILY SB1 WARNING ENTRY	0.051		
4182	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4183	X62162	B.burgdorferi gene for pC protein	0.41	<none></none>	<none></none>	<none></none>		

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		`	s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4184	Z81315	Human DNA sequence from fosmid F62D4 on chromosome 22q12-qter > :: emb Z81316 HSF 62D4A Human DNA sequence from fosmid F62D4 on chromosome 22, complete sequence	1.2	<none></none>	<none></none>	<none></none>	
4185	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
4186	L08108	Human low- affinity Fc- receptor IIB gene, exons 4-7.	0.0006	462387	IMMEDIATE- EARLY PROTEIN IE180 herpesvirus 1 (strain Kaplan) >gi 334071 (M34651) immediate-early protein [Pseudorabies virus]	0.25	
4187	AJ228330	Pinus pinaster reverse transcriptase gene of Line- retroelement (clone pPpLi1)	1.3	3108187	(AC004663) Notch 3 [Homo sapiens]	1.3	
4188	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4189	AF048991	Homo sapiens MutS homolog 5 (MSH5) gene, exons 13 through 25 and complete cds	0.002	3986756	[Mus musculus]	0.066	
4190	Z59608	H.sapiens CpG DNA, clone 165g8, reverse read cpg165g8.rt1a.	2e-014	1055183	(U40061) Similar to sodium-dependent phosphate transporter. [Caenorhabditis elegans]	2.4	

	<u> </u>	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4191	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
4192	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
4193	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	2128837	hypothetical protein MJ1401 - Methanococcus jannaschii >gi 1592049 (U67580) putative ATP dependent RNA helicase [Methanococcus jannaschii]	7.6	
4194	X99691	B.taurus DNA for agouti gene	9e-009	<none></none>	<none></none>	<none></none>	
4195	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	8e-008	306929	(M28696) IgG Fc receptor beta-Fc- gamma-RII [Homo sapiens]	0.64	
4196	U37521	Sus scrofa E- selectin gene, complete cds	0.042	539800	calcium-activated potassium channel mSlo - mouse >gi 347144 (L16912) mSlo [Mus musculus]	3.3	
4197	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds		<none></none>	<none></none>	<none></none>	
4198	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0002	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	· · · · · · · · · · · · · · · · · · ·	astN vs. Genbank)	•	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4199	V01087	Hemagglutinin gene of influenza virus strain A/duck/Ukraine/1/63 > :: gb J02109 FLAH AMU Influenza A/duck/ukraine/1/63 (h3n8), hemagglutinin (seg 4), cdna.	0.18	4038537	(AL021106) 1- evidence=predicte d by match; 1- match_accession= AA392988; 1- match_description =LD12167.5prime LD Drosophila melanogaster embryo BlueScript Drosophila melanogaster cDNA clone LD12167 5prime.; 1- match_species=Dr osop	8.5	
4200	X83107	H.sapiens Bmx mRNA for cytoplasmic tyrosine kinase	0.38	1147597	(U31221) viscerotropic leishmaniasis antigen [Leishmania tropica]	3.3	
4201	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.9	
4202	X71642	M.musculus GEG-154 mRNA	3.5	2760302	(D89074) hypothetical protein [Vibrio cholerae O139 fs1 phage]	1.5	
4203	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.7	
4204	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	1574918	(U19728) organic anion transporter [Raja erinacea]	5.8	
4205	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	<none></none>	<none></none>	<none></none>	

	<del></del>	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4206	U95094	Xenopus laevis   XL-INCENP   (XL-INCENP)   mRNA, complete   cds	4e-011	<none></none>	<none></none>	<none></none>	
4207	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>	
4208	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-011	<none></none>	<none></none>	<none></none>	
4209	U50523	Human BRCA2 region, mRNA sequence CG037	0	3121764	ARP2/3 COMPLEX 34 KD SUBUNIT	9e-026	
4210	X80909	H.sapiens alpha NAC mRNA	8e-050	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.9	
4211	AF039955	Homo sapiens liver CC chemokine-1 precursor	7e-006	<none></none>	<none></none>	<none></none>	
4212	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.6	
4213	L35670	Homo sapiens (subclone H8 10_g5 from P1 35 H5 C8) DNA sequence.	7e-017	<none></none>	<none></none>	<none></none>	
4214	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
4215	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4216	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4217	L33354	Lobostemon fruticosus Buek chloroplast trnL(UAA)- trnF(GAA) intergenic spacer DNA.	0.35	1483615	(Z77856) beta- glucosidase [Thermotoga neapolitana]	9	
			460		<u>.                                    </u>		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4218	Z12112	pWE15A cosmid	5e-033	987050	(X65335) lacZ	4e-008	
		vector DNA			gene product		
					[unidentified cloning vector]		
4219	X65279	pWE15 cosmid	2e-079	987050	(X65335) lacZ	3e-015	
	110027	vector DNA	20 0/2	,	gene product		
					[unidentified		
					cloning vector]		
4220	AF052165	Homo sapiens	e-170	2065177	(Y12790) Supt5h	1e-059	
		clone 24522 mRNA sequence			protein [Homo sapiens]		
4221	U95102	Xenopus laevis	2e-006	<none></none>	<none></none>	<none></none>	
	0,5102	mitotic	20 000	1,01,2	1,01,2		
		phosphoprotein					
		90 mRNA,					
4222	U95102	complete cds	2e-014	2072296	(U95098) mitotic	1.9	
4222	093102	Xenopus laevis mitotic	26-014	2072296	phosphoprotein 44	1.9	
		phosphoprotein			[Xenopus laevis]		
		90 mRNA,					
		complete cds					
4223	AF055024	Homo sapiens clone 24763	0	<none></none>	<none></none>	<none></none>	
		mRNA sequence					
4224	S39048	knob associated	0.39	<none></none>	<none></none>	<none></none>	
		histidine-rich					
		protein KAHRP					
4225	U95102	Xenopus laevis mitotic	4e-011	<none></none>	<none></none>	<none></none>	
		phosphoprotein					
		90 mRNA,					
		complete cds					
4226	U34377	Human tyrosine	2e-028	1709347	SERINE/THREO	8e-008	
		kinase TXK (txk) gene, exon 13.			NINE-PROTEIN KINASE NRK2		
		gene, exon 13.			(SERINE/THREO		
				:	NINE KINASE 2)		
					>gi 348245		
					(L20321) protein		
					serine/threonine kinase [Homo		
					sapiens]	]	
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		Nearest Neighbor			Nearest Neighbor			
95.0	`	astN vs. Genbank)	**************************************	(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4227	U25748	Pan troglodytes epididymal secretory protein precursor (EPI-1) mRNA, complete cds.	0	3182993	EPIDIDYMAL SECRETORY PROTEIN E1 PRECURSOR (EPI-1) (HE1) (EPIDIDYMAL SECRETORY PROTEIN 14.6) (ESP14.6) >gi 106343 pir  S2 5641 hypothetical protein - human >gi 2134519 pir  I5 3929 epididymal secretory protein 14.6 - crab-eating macaque human >gi 37477 (X676	7e-040		
4228	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4229	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.017	<none></none>	<none></none>	<none></none>		
4230	X74929	H.sapiens KRT8 mRNA for keratin 8	6e-036	<none></none>	<none></none>	<none></none>		
4231	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-007	<none></none>	<none></none>	<none></none>		
4232	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6		
4233	U41010	Caenorhabditis elegans cosmid T05A12	4.2	<none></none>	<none></none>	<none></none>		
4234	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-007	1363925	hypothetical protein 2 - North American opossum (fragment) >gi 897721 (Z48955) ORF-2, putative RT [Didelphis	4.7		

	1	Nearest Neighbor		Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					virginiana]		
4235	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	439493	(D26086) zinc- finger protein [Petunia x hybrida]	8.5	
4236	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	2501599	HYPOTHETICAL 29.1 KD PROTEIN W06E11.4 IN CHROMOSOME III >gi 669022 (U20862) W06E11.4 gene product [Caenorhabditis elegans]	0.002	
4237	X94118	P.falciparum PK4 gene	1.2	<none></none>	<none></none>	<none></none>	
4238	Z18944	S.cerevisiae BDF1 gene	7.30E-01	2119161	unknown - chicken (fragment) >gi 537433	0.61	
4239	AF031939	Mus musculus RalBP1- associated EH domain protein Reps1 (reps1) mRNA, complete cds	e-154	2677843	(AF031939) RalBP1-associated EH domain protein Reps1	5e-016	
4240	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.9	
4241	L35566	Gallus gallus homeobox protein (LH-2) mRNA, complete cds.	3e-044	1708809	HOMEOBOX PROTEIN LH-2 >gi 508712	4e-021	
4242	Z83086	H.sapiens Fanconi anaemia group A gene, exon 29	3.00E-07	<none></none>	<none></none>	<none></none>	

	1	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4243	U63810	Homo sapiens WD40 protein Ciao 1 mRNA, complete cds	0.00E+00	3219331	(AC004020) Unknown gene product [Homo sapiens]	1e-096	
4244	U15110	Mycoplasma capricolum ptsI- crr operon phosphocarrier protein enzyme I (ptsI) and phosphocarrier protein enzyme IIA (crr) genes, complete cds, and lipopolysaccharid e biosynthesis (kdtB) gene, complete cds.	1.1	<none></none>	<none></none>	<none></none>	
4245	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4246	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-011	730888	OCTAPEPTIDE- REPEAT PROTEIN T2	1.4	
4247	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.074	<none></none>	<none></none>	<none></none>	
4248	AJ224152	Plasmodium berghei gene encoding cdc2- related kinase 2	0.54	<none></none>	<none></none>	<none></none>	
4249	M24971	D.discoideum CT-rich satellite rDNA, clone pCT11.	2e-008	119110	EBNA-1 NUCLEAR PROTEIN herpesvirus 4 (strain B95-8) >gi 1334880 (V01555) BKRF1 encodes EBNA-1 protein, latent cycle gene. [Human herpesvirus 4]	2e-009	
4250	Z72969	S.cerevisiae chromosome VII reading frame ORF YGR184c	1.2	<none></none>	<none></none>	<none></none>	

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	Nearest Neighbor			Nearest Neighbor			
	•	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4251	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4252	AJ224326	Homo sapiens mRNA for putative ribulose- 5-phosphate- epimerase, partial cds	0	<none></none>	<none></none>	<none></none>	
4253	U45245	Homo sapiens paired-box protein PAX2 (PAX2) gene, promoter and exon 1	2.1	<none></none>	<none></none>	<none></none>	
4254	AE001157	Borrelia burgdorferi (section 43 of 70) of the complete genome	0.63	<none></none>	<none></none>	<none></none>	
4255	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.8	
4256	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0005	2773162	(AF039595) sulfonylurea receptor 1B [Rattus norvegicus]	9.6	
4257	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5e-009	<none></none>	<none></none>	<none></none>	
4258	L11130	Influenza A/gull/MD/19/77 (H2N8) hemagglutinin	0.67	<none></none>	<none></none>	<none></none>	
4259	AB018270	Homo sapiens mRNA for KIAA0727 protein, partial cds	0	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3	
4260	U67494	Methanococcus jannaschii section 36 of 150 of the complete genome	0.014	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)  ACCESSION   DESCRIPTION   P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION			
4261	L09209	Homo sapiens amyloid protein homologue mRNA, complete cds > :: gb I13782 I13782 Sequence 12 from patent US 5441931 > :: gb I68752 I68752 Sequence 12 from patent US 5677146	6e-089	<none></none>	<none></none>	<none></none>	
4262	M27866	Human retinoblastoma susceptibility protein gene, exon 27. > :: gb I09392  Sequence 25 from Patent WO 8906703	e-158	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.7	
4263	U59629	Human transcription factor LZIP-alpha mRNA, complete cds	1e-052	2828799	(U55386) unknown [Anabaena PCC7120]	0.097	
4264	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	3176395	(AB015041) PIF1 [Caenorhabditis elegans]	3e-005	
4265	AF069250	Homo sapiens okadaic acid- inducible phosphoprotein (OA48-18) mRNA, complete cds	2e-068	3037018	(AF041330) NADH dehydrogenase subunit 5 [Bodo saltans]	0.002	
4266	M11560	Human aldolase A mRNA, complete cds.	0.00E+00	113606	FRUCTOSE- BISPHOSPHATE ALDOLASE A fructose- bisphosphate aldolase (EC 4.1.2.13) A - human sapiens]	5e-055	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4267	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-009	<none></none>	<none></none>	<none></none>	
4268	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-005	2688708	(AE001176) conserved hypothetical protein [Borrelia burgdorferi]	8.5	
4269	L35566	Gallus gallus homeobox protein (LH-2) mRNA, complete cds.	6e-041	1708809	HOMEOBOX PROTEIN LH-2 >gi 508712	7e-019	
4270	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1.00E-11	1709997	DNA REPAIR PROTEIN RAD18 >gi 1150622 protein rad18 [Schizosaccharom yces pombe]	6e-027	
4271	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-009	586442	NUCLEOPORIN NUP170 (NUCLEAR PORE PROTEIN NUP170) >gi 626192 pir  S4 5429 probable membrane protein YBL079w - yeast (Saccharomyces cerevisiae) cerevisiae] >gi 536127 (Z35840) ORF YBL079w	0.44	
4272	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-013	<none></none>	<none></none>	<none></none>	
4273	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4274	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor			Nearest Neighbor			
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SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4275	X00334	Drosophila virilis simple DNA sequence (pDv- 19)	6e-010	119110	EBNA-1 NUCLEAR PROTEIN herpesvirus 4 (strain B95-8) >gi 1334880 (V01555) BKRF1 encodes EBNA-1 protein, latent cycle gene. [Human herpesvirus 4]	2e-016		
4276	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>		
4277	AF069250	Homo sapiens okadaic acidinducible phosphoprotein (OA48-18) mRNA, complete cds	2e-068	3037018	(AF041330) NADH dehydrogenase subunit 5 [Bodo saltans]	0.002		
4278	Y10183	H.sapiens mRNA for MEMD protein	e-162	<none></none>	<none></none>	<none></none>		
4279	D86960	Human mRNA for KIAA0205 gene, complete cds	2e-078	<none></none>	<none></none>	<none></none>		
4280	X65319	Cloning vector pCAT-Enhancer	3e-081	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015		
4281	X86693	H.sapiens mRNA for hevin like protein	0.18	<none></none>	<none></none>	<none></none>		
4282	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0001	<none></none>	<none></none>	<none></none>		
4283	M33156	A.aegypti D7 gene, exons 1-5.	1.30E+00	<none></none>	<none></none>	<none></none>		
4284	X02317	Human mRNA for Cu/Zn superoxide dismutase (SOD)	0	218564	(D90358) HB- SOD [Schizosaccharom yces pombe]	7e-032		

		Nearest Neighbor			Nearest Neighbor			
	` `	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4285	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-04	<none></none>	<none></none>	<none></none>		
4286	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-04	<none></none>	<none></none>	<none></none>		
4287	X02317	Human mRNA for Cu/Zn superoxide dismutase (SOD)	0	134611	SUPEROXIDE DISMUTASE (CU-ZN) dismutase (aa 1- 154) [Homo sapiens] >gi 338276 (K00065) superoxide dismutase [Homo sapiens] >gi 1237407 (L44139) Cu/Zn- superoxide dismutase [Homo sapiens]	2e-079		
4288	X04408	Human mRNA for coupling protein G(s) alpha subunit adenylyl cyclase)	0	386748	(M14631) guanine nucleotide-binding protein alpha subunit	2e-073		
4289	M28161	Rabbit MHC class II RLA-DR- alpha gene, complete cds.	2.4	<none></none>	<none></none>	<none></none>		
4290	U33956	Human Down Syndrome region of chromosome 21, genomic sequence, clone A12H1-1F8.	0.37	<none></none>	<none></none>	<none></none>		
4291	U90331	Mus musculus neural plakophilin related arm- repeat protein (NPRAP) mRNA, complete cds	0.15	135063	SUPPRESSOR OF SABLE PROTEIN fruit fly (Drosophila melanogaster) >gi 158517 (M57889) su(s) protein	5.2		

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	PVALUE	
ID							
					[Drosophila		
					melanogaster]		
4292	AF045531	Homo sapiens	0.005	<none></none>	<none></none>	<none></none>	
		germline				1,01,2	
		chromosome 22,					
		22q11.2 region					
4293	D86960	Human mRNA	2e-078	<none></none>	<none></none>	<none></none>	
		for KIAA0205					
		gene, complete					
1204	U95094	cds	2e-005	Aloves	AIONES	<none></none>	
4294	U93094	Xenopus laevis XL-INCENP	2e-005	<none></none>	<none></none>	<nune></nune>	
		(XL-INCENP)					
		mRNA, complete					
		cds					
4295	U95098	Xenopus laevis	5.00E-03	<none></none>	<none></none>	<none></none>	
		mitotic					
		phosphoprotein					
		44 mRNA, partial					
4296	U17073	cds Neurospora	0.041	3152938	(AF065482)	0.83	
4290	017073	crassa frequency	0.041	3132936	sorting nexin 2	0.63	
		(frq) mRNA,			[Homo sapiens]		
		complete cds.			[		
4297	M93051	Pisum sativum	0.2	<none></none>	<none></none>	<none></none>	
		ascorbate					
		peroxidase					
		(ApxI) gene,					
4200	1100153	complete cds.	0.27	AIONES	AIONE:	AIONE	
4298	U28153	Caenorhabditis elegans UNC-76	0.37	<none></none>	<none></none>	<none></none>	
		(unc-76) gene,					
		complete cds.					
4299	U28153	Caenorhabditis	0.37	<none></none>	<none></none>	<none></none>	
		elegans UNC-76					
		(unc-76) gene,					
		complete cds.					

 $(a_{n},a_{n},\ldots,a_{n}) = (a_{n},\ldots,a_{n}) = (a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots,a_{n},\ldots$ 

-51	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4300	U20240	Human C/EBP gamma mRNA, complete cds > :: gb G28590 G285 90 human STS SHGC-35371.	e-141	1705750	CCAAT/ENHAN CER BINDING PROTEIN GAMMA (C/EBP GAMMA) >gi 1363931 pir  J C4243 transcription CCAAT enhancer binding protein- gamma - human >gi 727294 (U20240) C/EBP gamma [Homo sapiens]	1e-011	
4301	Y16359	Calonectris diomedea random amplified polymorphic DNA, clone Cd- O8fl	4e-075	595780	(U13871) lacZ alpha peptide [Cloning vector]	0.0001	
4302	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-013	<none></none>	<none></none>	<none></none>	
4303	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4304	U90331	Mus musculus neural plakophilin related arm- repeat protein (NPRAP) mRNA, complete cds	0.15	135063	SUPPRESSOR OF SABLE PROTEIN fruit fly (Drosophila melanogaster) >gi 158517 (M57889) su(s) protein [Drosophila melanogaster]	5.2	
4305	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4306	D86960	Human mRNA for KIAA0205 gene, complete cds	0	1653865	(D90917) UDP-N-acetylglucosamine -N-acetylmuramyl- (pentape ptide) pyrophosphoryl - undecaprenol N-acetylglucosamine transferase [Synechocystis sp.]	4.40E+00	

	Nearest Neighbor			Nearest Neighbor			
	`	lastN vs. Genbank)	I	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4307	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-013	4105520	(AF046933) carboxysome structural polypeptide	2.4	
4308	Y14723	Choanomphalus incertus mitochondrial cytochrome c oxidase subunit I gene, partial	0.36	<none></none>	<none></none>	<none></none>	
4309	AB018327	Homo sapiens mRNA for KIAA0784 protein, partial cds	0	3882289	(AB018327) KIAA0784 protein [Homo sapiens]	4e-041	
4310	AB007860	Homo sapiens KIAA0400 mRNA, complete cds	0	<none></none>	<none></none>	<none></none>	
4311	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4312	U96440	Drosophila melanogaster cut gene, partial sequence	0.053	119110	EBNA-1 NUCLEAR PROTEIN herpesvirus 4 (strain B95-8) >gi 1334880 (V01555) BKRF1 encodes EBNA-1 protein, latent cycle gene. [Human herpesvirus 4]	0.0004	
4313	X64707	H.sapiens BBC1 mRNA	3e-090	1350662	60S RIBOSOMAL PROTEIN L13 (A52)	2e-025	
4314	U67522	Methanococcus jannaschii section 64 of 150 of the complete genome	0.38	<none></none>	<none></none>	<none></none>	
4315	M11560	Human aldolase A mRNA, complete cds.	0.00E+00	113606	FRUCTOSE- BISPHOSPHATE ALDOLASE A fructose- bisphosphate aldolase (EC 4.1.2.13) A - human sapiens]	5e-055	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4316	X92098	H.sapiens mRNA for transmembrane protein rnp24	e-123	3914237	COP-COATED VESICLE MEMBRANE PROTEIN P24 PRECURSOR (P24A) (RNP24) >gi 1212965 gnl PI D e205529	1e-017	
4317	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4318	D86960	Human mRNA for KIAA0205 gene, complete cds	0	1653865	(D90917) UDP-N-acetylglucosamine -N-acetylmuramyl- (pentape ptide) pyrophosphoryl - undecaprenol N- acetylglucosamine transferase [Synechocystis sp.]	4.40E+00	
4319	M83094	Homo sapiens cytosolic selenium-dependent glutathione peroxidase gene, complete cds, and rhoh12 gene, 3' end.	0.00E+00	<none></none>	<none></none>	<none></none>	
4320	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5e-015	<none></none>	<none></none>	<none></none>	
4321	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
4322	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	3.3	<none></none>	<none></none>	<none></none>	
4323	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1.40E-02	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4324	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4325	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	<none></none>	<none></none>	<none></none>	
4326	AF088034	Homo sapiens full length insert cDNA clone ZC24F03	0	854598	(X87611) ORF YJR83.18 [Saccharomyces cerevisiae]	2e-024	
4327	U47322	Cloning vector DNA, complete sequence.	6.00E-06	<none></none>	<none></none>	<none></none>	
4328	U47322	Cloning vector DNA, complete sequence.	6.00E-06	<none></none>	<none></none>	<none></none>	
4329	D86960	Human mRNA for KIAA0205 gene, complete cds	0.00E+00	1653865	(D90917) UDP-N-acetylglucosamine -N-acetylmuramyl- (pentape ptide) pyrophosphoryl - undecaprenol N- acetylglucosamine transferase [Synechocystis sp.]	1.4	
4330	Z70316	D.melanogaster mRNA for tyramine-beta- hydroxylase	1.5	<none></none>	<none></none>	<none></none>	
4331	L28010	Homo sapiens HnRNP F protein mRNA, complete cds	3e-070	1710628	HETEROGENEO US NUCLEAR RIBONUCLEOPR OTEIN F (HNRNP F) >gi 631210 pir  S4 3484 hnRNP F protein - human >gi 452048 (L28010) HnRNP F protein [Homo sapiens]	2e-005	
4332	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1.40E-01	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor		Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4333	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.13	<none></none>	<none></none>	<none></none>	
4334	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.004	1723286	VERY HYPOTHETICAL 11.9 KD PROTEIN C4H3.12C IN CHROMOSOME I >gi 1184025 (Z69380) unknown	3.1	
4335	<none></none>	<none></none>	<none></none>	2314752	(AE000654) rare lipoprotein A (rlpA) [Helicobacter pylori]	7.3	
4336	AB007963	Homo sapiens mRNA for KIAA0494 protein, complete cds	8e-078	3413938	(AB007963) KIAA0494 protein [Homo sapiens]	1.00E-11	
4337	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4338	X12597	Human mRNA for high mobility group-1 protein	3e-048	123371	HIGH MOBILITY GROUP PROTEIN HMG1 protein HMG-1 - pig >gi 164490 (M21683) non- histone protein HMG1 [Sus scrofa]	0.006	
4339	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-013	2853095	(AL021767) very hypothetical protein	0.043	
4340	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-007	<none></none>	<none></none>	<none></none>	
4341	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-006	3063453	(AC003981) F22O13.15 [Arabidopsis thaliana]	4.5	

		Nearest Neighbor			Nearest Neighbor			
	· · · · · · · · · · · · · · · · · · ·	astN vs. Genbank)			s. Non-Redundant Pr			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4342	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4.00E-11	231629	BILE-SALT- ACTIVATED LIPASE PRECURSOR ESTER LIPASE) (STEROL ESTERASE) (CHOLESTEROL ESTERASE) salt- activated lipase [Homo sapiens] sapiens]	9.6		
4343	L31732	Human STS UT643, 5' primer bind.	1.6	<none></none>	<none></none>	<none></none>		
4344	AF037332	Homo sapiens Eph-like receptor tyrosine kinase hEphB1b (EphB1) mRNA, complete cds	0.66	<none></none>	<none></none>	<none></none>		
4345	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4346	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2.00E-05	<none></none>	<none></none>	<none></none>		
4347	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-007	<none></none>	<none></none>	<none></none>		
4348	Z30961	H.sapiens DNA for Mhc Alu elements	7.00E-17	728835	!!!! ALU SUBFAMILY SC WARNING ENTRY	0.5		
4349	U34887	Yeast integrating vector pRS306 containing a fragment of lacZ.	7e-068	3152967	(Y14016) hypothetical protein	9		
4350	D28124	Human mRNA for unknown product, complete cds	0	1825638	(U88172) similar to protein-tyrosine phosphatase	0.062		
4351	AF069503	Carcharhinus plumbeus microsatellite repeat region	4.20E+00	<none></none>	<none></none>	<none></none>		

	N	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4352	AF069503	Carcharhinus plumbeus microsatellite repeat region	4.20E+00	<none></none>	<none></none>	<none></none>	
4353	D10848	Alkalophilic Bacillus sp. genomic DNA for lipo-penicillinase	0.033	<none></none>	<none></none>	<none></none>	
4354	D28124	Human mRNA for unknown product, complete cds	0	1825638	(U88172) similar to protein-tyrosine phosphatase	0.062	
4355	U19482	Mus musculus C10-like chemokine mRNA, complete cds	3.70E+00	<none></none>	<none></none>	<none></none>	
4356	AF050068	Homo sapiens growth arrest specific 11	1.4	1916844	(U82987) Bcl-2 binding component 3 [Homo sapiens]	0.042	
4357	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
4358	AE000026	Mycoplasma pneumoniae section 26 of 63 of the complete genome	1.3	<none></none>	<none></none>	<none></none>	
4359	<none></none>	<none></none>	<none></none>	2114321	(D88733) membrane glycoprotein [Equine herpesvirus 1]	8.00E-01	
4360	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

	4	Nearest Neighbor	•	Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID`							
4361	Y07660	M.tuberculosis accBC gene	2e-068	465847	HYPOTHETICAL 66.5 KD PROTEIN F02A9.5 IN CHROMOSOME III >gi 280542 pir  S2 8313 hypothetical protein F02A9.5 - Caenorhabditis elegans Genefinder; similar to Propionyl-CoA carboxylase beta chain; cDNA EST EMBL:M89018 comes from this gene; cDNA EST EMBL:D2806	4e-079	
4362	U12022	Human calmodulin (CALM1) gene, exons 2,3,4,5 and 6, and complete cds	e-127	<none></none>	<none></none>	<none></none>	
4363	AC001178	Homo sapiens (subclone 2_g12 from BAC H94) DNA sequence	3.00E-28	<none></none>	<none></none>	<none></none>	
4364	<none></none>	<none></none>	<none></none>	4063042	(AF068065) GP900; mucin-like glycoprotein [Cryptosporidium parvum]	0.52	
4365	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4366	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1.00E-12	<none></none>	<none></none>	<none></none>	
4367	X14448	Human GLA gene for alpha-D- galactosidase A (EC 3.2.1.22)	3	<none></none>	<none></none>	<none></none>	

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	Nearest Neighbor			Nearest Neighbor			
GD0		astN vs. Genbank)	DAVALLED	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4368	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	5.00E-04	3873753	(Z66519) similar to phytoene synthase precursor; cDNA EST yk340f7.3 comes from this gene; cDNA EST yk340f7.5 comes from this gene [Caenorhabditis elegans]	2e-008	
4369	X04098	Human mRNA for cytoskeletal gamma-actin	0	<none></none>	<none></none>	<none></none>	
4370	M13452	Human lamin A mRNA, 3'end.	0	125962	LAMIN A (70 KD LAMIN)	3e-057	
4371	AF068863	Homo sapiens oligodendrocyte-specific protein	3.4	<none></none>	<none></none>	<none></none>	
4372	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1.40E-01	<none></none>	<none></none>	<none></none>	
4373	L04636	Homo sapiens pre-mRNA splicing factor 2 p32 subunit (SF2p32) mRNA, complete cds.	0	730772	COMPLEMENT COMPONENT 1, Q SUBCOMPONEN T BINDING PROTEIN PRECURSOR (GLYCOPROTEI N GC1QBP) (GC1Q-R PROTEIN) (HYALURONAN -BINDING PROTEIN 1) chain precursor - human >gi 338045 (L04636) splicing factor [Homo sapiens] >gi 472956 (X75913) gCIq-R [Homo sapiens] >gi	2e-050	
4374	M59832	Human merosin mRNA, 3' end.	0.043	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4375	<none></none>	<none></none>	<none></none>	188864	(M74027) mucin [Homo sapiens]	0.042	
4376	X17206	Human mRNA for LLRep3	0	88570	ribosomal protein S2 - human (fragment) sapiens]	6e-078	
4377	X17206	Human mRNA for LLRep3	0	88570	ribosomal protein S2 - human (fragment) sapiens]	6e-078	
4378	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4379	X98420	S.shibatae topR gene	1.10E+00	2746890	(AF040655) No definition line found [Caenorhabditis elegans]	9.3	
4380	X98420	S.shibatae topR gene	1.10E+00	2746890	(AF040655) No definition line found [Caenorhabditis elegans]	9.3	
4381	X75787	P.falciparum (FAF-2) mRNA for aspartic hemoglobinase	4	<none></none>	<none></none>	<none></none>	
4382	AF044209	Homo sapiens nuclear receptor co-repressor N- CoR mRNA, complete cds	0	3510603	(AF044209) nuclear receptor co-repressor N- CoR [Homo sapiens]	4e-029	
4383	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4384	X64707	H.sapiens BBC1 mRNA	e-110	1350662	60S RIBOSOMAL PROTEIN L13 (A52)	0.003	
4385	Z70316	D.melanogaster mRNA for tyramine-beta- hydroxylase	1.5	<none></none>	<none></none>	<none></none>	
4386	AF000371	Vitis vinifera UDP glucose:flavonoid 3-o- glucosyltransferas e mRNA, partial cds	0.19	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor			Nearest Neighbor			
000	,	astN vs. Genbank)	I DAVALLED	(BlastX vs. Non-Redundant Proteins)  ACCESSION   DESCRIPTION   P VALUE				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION				
4387	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.8		
4388	AE000688	Aquifex aeolicus section 20 of 109 of the complete genome	3.8	<none></none>	<none></none>	<none></none>		
4389	L05612	Dictyostelium purpureum DNA sequence, repeat region.	2.8	<none></none>	<none></none>	<none></none>		
4390	U33761	Human cyclin A/CDK2- associated p45 (Skp2) mRNA, complete cds	2e-079	2134952	cyclin A/CDK2- associated p45 - human sapiens]	1e-025		
4391	U48288	Rattus norvegicus A-kinase anchoring protein AKAP 220 mRNA, complete cds	0.48	<none></none>	<none></none>	<none></none>		
4392	AB007963	Homo sapiens mRNA for KIAA0494 protein, complete cds	0.00E+00	3413938	(AB007963) KIAA0494 protein [Homo sapiens]	6e-071		
4393	<none></none>	<none></none>	<none></none>	119110	EBNA-1 NUCLEAR PROTEIN herpesvirus 4 (strain B95-8) >gi 1334880 (V01555) BKRF1 encodes EBNA-1 protein, latent cycle gene. [Human herpesvirus 4]	6e-027		
4394	U527 <b>8</b> 4	Ansonia muelleri CMNH H1476 16S rRNA gene, mitochondrial gene encoding mitochondrial rRNA, partial sequence	0.014	<none></none>	<none></none>	<none></none>		

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
4395	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4396	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4397	U38376	Rattus norvegicus cytosolic phospholipase A2 mRNA, complete cds	1.1	<none></none>	<none></none>	<none></none>	
4398	U78770	Mus musculus spasmolytic polypeptide (mSP) gene, complete cds	0.028	<none></none>	<none></none>	<none></none>	
4399	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.12	<none></none>	<none></none>	<none></none>	
4400	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-014	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.8	
4401	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0003	<none></none>	<none></none>	<none></none>	
4402	X70288	H.sapiens gene for thioredoxin, exons 4 and 5	3e-030	<none></none>	<none></none>	<none></none>	
4403	X76683	Plasmid vector pHM2 betalactamase gene	7e-080	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	
4404	X69295	H.sapiens MSX2 mRNA for transcription factor	0.43	<none></none>	<none></none>	<none></none>	
4405	U20371	Mus musculus homeobox protein (Hoxa11) gene, complete cds.	0.6	<none></none>	<none></none>	<none></none>	

	7	Nearest Neighbor		Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4406	D49842	Rabbit mRNA for CD86, complete cds	1.10E+00	135554	TETRACYCLINE RESISTANCE PROTEIN Bacillus cereus plasmid pBC16 >gi 72838 pir  YTS OG tetracycline resistance protein- Streptococcus agalactiae plasmid pMV158 >gi 80428 pir  JQ1 211 tetracycline resistance protein- Bacillus sp. plasmid pTB19 >gi 151696 (M63	1.4	
4407	AB007194	Oryza sativa mRNA for fructose-1,6- bisphosphatase (plastidic isoform), complete cds	3.5	<none></none>	<none></none>	<none></none>	
4408	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1e-007	<none></none>	<none></none>	<none></none>	
4409	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4410	U28924	Pisum sativum cytosolic glutamine synthetase	0.008	3769486	(AF074946) DNA polymerase [hemorrhagic enteritis virus]	1.3	
4411	D30783	Homo sapiens mRNA for epiregulin, complete cds	0	1723438	HYPOTHETICAL 52.3 KD PROTEIN C56F8.06C IN CHROMOSOME I PRECURSOR >gi 1204228 (Z69728) unknown [Schizosaccharom yces pombe]	0.13	
4412	AJ012449	Homo sapiens mRNA for NS1- binding protein	0	3851214	(AJ012449) NS1- binding protein [Homo sapiens]	4e-088	

		Nearest Neighbor			Nearest Neighbor			
0.00		lastN vs. Genbank)	I	(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4413	X62357	H.sapiens Alu repeat (clones 2- 48)	1e-006	<none></none>	<none></none>	<none></none>		
4414	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6.00E-05	<none></none>	<none></none>	<none></none>		
4415	Z15015	D.pulex mitochondrion genes for NADH dehydrogenase subunit 2, cytochrome C oxidase subunit I, tRNA-Val, tRNA-Ile, tRNA-Gln, tRNA-fMet, tRNA-Trp, tRNA-Cys, tRNA-Tyr, small subunit rRNA, large subunit rRNA	2.2	1076802	extensin-like protein - maize >gi 600118	8e-027		
4416	D87942	Homo sapiens mRNA for alpha(1,2)fucosyl transferase, complete cds	2e-027	728838	!!!! ALU SUBFAMILY SX WARNING ENTRY	7.5		
4417	D86977	Human mRNA for KIAA0224 gene, complete cds	0	3024898	PUTATIVE PRE-MRNA SPLICING FACTOR ATP-DEPENDENT RNA HELICASE KIAA0224 (HA4657) putative ATP-dependent RNA helicase K03H1.2 of C.elegans(S41025) [Homo sapiens] >gi 3123906 (AF038391) pre- mRNA splicing factor [Homo sapiens]	2e-053		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4418	L28010	Homo sapiens HnRNP F protein mRNA, complete cds	0	1710628	HETEROGENEO US NUCLEAR RIBONUCLEOPR OTEIN F (HNRNP F) >gi 631210 pir  S4 3484 hnRNP F protein - human >gi 452048 (L28010) HnRNP F protein [Homo sapiens]	5e-045	
4419	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
4420	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
4421	X14313	Arabidopsis CRB gene for 12S seed storage protein > gene, exons 1-4.	0.24	<none></none>	<none></none>	<none></none>	
4422	X14313	Arabidopsis CRB gene for 12S seed storage protein > gene, exons 1-4.	0.24	<none></none>	<none></none>	<none></none>	
4423	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
4424	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	3.00E-08	1125753	(U42833) coded for by C. elegans cDNA CEESN37F; Similar to ammonium transport protein. [Caenorhabditis elegans]	1e-019	
4425	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-05	<none></none>	<none></none>	<none></none>	

	N	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
4426	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	1125753	(U42833) coded for by C. elegans cDNA CEESN37F; Similar to ammonium transport protein. [Caenorhabditis elegans]	2e-008	
4427	AF053649	Homo sapiens cellular apoptosis susceptibility protein (CSE1) gene, exons 15 and 16	3e-008	<none></none>	<none></none>	<none></none>	
4428	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1e-007	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	8.8	
4429	X94253	S.scrofa mRNA for heterogeneous nuclear ribonucleoprotein	6e-023	<none></none>	<none></none>	<none></none>	
4430	AF005039	Homo sapiens secretory carrier membrane protein	0	2232243	(AF005039) secretory carrier membrane protein [Homo sapiens]	8e-008	
4431	AF037332	Homo sapiens Eph-like receptor tyrosine kinase hEphB1b (EphB1) mRNA, complete cds	0.12	3861156	(AJ235272) unknown [Rickettsia prowazekii]	0.37	
4432	D28124	Human mRNA for unknown product, complete cds	7e-067	<none></none>	<none></none>	<none></none>	
4433	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-013	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4434	M93426	Human protein tyrosine phosphatase zetapolypeptide (PTPRZ) mRNA, complete cds. > :: gb G20044 G200 44 sWSS1987 Eric D. Green Homo sapiens STS genomic, sequence tagged site [Homo sapiens]	0	400199	PROTEIN- TYROSINE PHOSPHATASE ZETA PRECURSOR (R- PTP-ZETA) >gi 476869 pir  A4 6151 protein- tyrosine- phosphatase (EC 3.1.3.48), receptor type zeta - human >gi 190744 (M93426) protein tyrosine phosphatase zeta- polypeptide [Homo sapiens]	4e-051	
4435	Ū54562	Human translation initiation factor eIF3 p48 subunit (Int-6) mRNA, complete cds	0	2498490	VIRAL INTEGRATION SITE PROTEIN INT-6 >gi 1854579 (L35556) Int-6 [Mus musculus] sapiens] >gi 2351382 (U54562) eIF3- p48 [Homo sapiens] sapiens]	e-110	
4436	U54562	Human translation initiation factor eIF3 p48 subunit (Int-6) mRNA, complete cds	0	2498490	VIRAL INTEGRATION SITE PROTEIN INT-6 >gi 1854579 (L35556) Int-6 [Mus musculus] sapiens] >gi 2351382 (U54562) eIF3- p48 [Homo sapiens] sapiens]	e-110	
4437	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.12	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4438	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-009	<none></none>	<none></none>	<none></none>	
4439	X03558	Human mRNA for elongation factor 1 alpha subunit	0	1169475	ELONGATION FACTOR 1- ALPHA 1	4e-083	
4440	J03607	Human 40-kDa keratin intermediate filament precursor gene.	0	1070608	keratin 19, type I, cytoskeletal - human sapiens]	4e-058	
4441	<none></none>	<none></none>	<none></none>	4063042	(AF068065) GP900; mucin-like glycoprotein [Cryptosporidium parvum]	0.011	
4442	<none></none>	<none></none>	<none></none>	4063042	(AF068065) GP900; mucin-like glycoprotein [Cryptosporidium parvum]	0.011	
4443	Y13401	Homo sapiens CD3 delta gene, enhancer sequence	8e-008	<none></none>	<none></none>	<none></none>	
4444	X04409	Human mRNA for coupling protein G(s) alpha-subunit (alpha-S1) (stimulatory regulatory component Gs of adenylyl cyclase)	0	71879	GTP-binding regulatory protein Gs alpha chain G- s-alpha-4 [Homo sapiens]	7e-092	
4445	AF038958	Homo sapiens synaptic glycoprotein SC2 spliced variant mRNA, complete cds	1e-072	3329386	(AF038958) synaptic glycoprotein SC2 spliced variant	6e-019	
4446	D17244	Human HepG2 3' region MboI cDNA, clone hmd4h04m3	1e-075	2500256	50S RIBOSOMAL PROTEIN L13 protein L13 [Streptomyces coelicolor]	0.043	

	N	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4447	<none></none>	<none></none>	<none></none>	4063042	(AF068065) GP900; mucin-like glycoprotein [Cryptosporidium parvum]	0.005		
4448	M24597	Beet curly top virus (clone pBCT028) DNA, complete genome.	4.1	<none></none>	<none></none>	<none></none>		
4449	U59706	Gallus gallus alternatively spliced AMPA glutamate receptor, isoform GluR2 flop, (GluR2) mRNA, partial cds.	0.014	3283975	(AF072521) poly- (ADPribosyl)- transferase homolog PARP	0.02		
4450	AJ010014	Homo sapiens mRNA for M96A protein	0	3342452	(AF072814) PHD finger DNA binding protein isoform 1	2e-029		
4451	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4452	X06960	Aspergillus nidulans mitochondrial DNA for cytochrome oxidase subunit 3, tRNA-Tyr	0.23	<none></none>	<none></none>	<none></none>		
4453	L01089	Human profilaggrin (FLG) gene exons 2-3, 5'end.		<none></none>	<none></none>	<none></none>		
4454	X65319	Cloning vector pCAT-Enhancer	1e-071	987050	(X65335) lacZ gene product [unidentified cloning vector]	1e-014		
4455	X87212	H.sapiens mRNA for cathepsin C	0	1582221	prepro-cathepsin C [Homo sapiens]	6e-046		
4456	X53123	Cloning vector pAST 19a for C. elegans	5	<none></none>	<none></none>	<none></none>		
4457	D15057	Human mRNA for DAD-1, complete cds	0	2944452	(AF051310) defender against death 1 [Mus musculus]	1e-015		

and the same	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4458	X83860	H.sapiens mRNA for prostaglandin E receptor (EP3c)	1.2	2137044	unknown protein - rabbit (fragment) cuniculus]	7e-014	
4459	M95058	Rattus rattus steroid 5-alpha- reductase 2 mRNA, complete cds.	0.42	<none></none>	<none></none>	<none></none>	
4460	AF044588	Homo sapiens protein regulating cytokinesis 1	2e-043	2865521	(AF044588) protein regulating cytokinesis 1; PRC1 [Homo sapiens]	4e-015	
4461	X54282	Human chromosome 11 DNA, approx. 20 kb 3' of beta- globin gene, nuclear scaffold associated region	0.014	1911867	cadherin 3 [Caenorhabditis elegans, Peptide, 3337 aa]	9.8	
4462	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	3875640	(Z92781) F09C3.3 [Caenorhabditis elegans]	9.6	
4463	M73791	Human novel gene mRNA, complete cds.	0	1172810	60S RIBOSOMAL PROTEIN L10 (QM PROTEIN HOMOLOG) >gi 543339 pir  JC 2013 ribosomal protein L10, cytosolic - mouse >gi 2143959 pir  J C4911 ribosomal protein L10 - rat >gi 407466 (X75312) QM protein [Mus musculus] >gi 410742 (M93980) 24.6 kda protein [Mus musc	7e-085	
4464	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor			Nearest Neighbor			
OF C	```	lastN vs. Genbank)  DESCRIPTION	DVALUE		s. Non-Redundant Pr			
SEQ ID	ACCESSION		P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4465	Z27116	S.cerevisiae HBS1, MRP-L20 and PRP-16 genes	0.058	<none></none>	<none></none>	<none></none>		
4466	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4467	M96575	Drosophila melanogaster collagen type IV gene, complete cds.	3.60E+00	<none></none>	<none></none>	<none></none>		
4468	D50010	Human DNA for alpha-platelet- derived growth factor receptor, exon 15	1e-006	<none></none>	<none></none>	<none></none>		
4469	X70649	Homo sapiens DDX1 gene, complete CDS	0	539572	DEAD box protein RB - human	3e-036		
4470	AJ223377	Puumala virus S- segment RNA	1.4	<none></none>	<none></none>	<none></none>		
4471	Y14599	Staphylococcus xylosus lacR, lacP, lacH genes and 2 ORF's	1.4	3659505	(AC005084) similar to mouse mCASK-A; similar to e1288039	0.63		
4472	X13336	Spinach plastid genes rps3, rps19, rpl14, rpl16 and rpl22 for ribosomal proteins S3, S19, L14, L16 and L22	0.15	1330375	(U58758) similar to rat GAP- associated protein p190	0.27		
4473	AF056022	Homo sapiens p60 katanin mRNA, complete cds	0	3283072	(AF056022) p60 katanin [Homo sapiens]	7e-029		
4474	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>		
4475	M86849	Human connexin 26 (GJB2) mRNA.	0	127542	ALDOSE 1- EPIMERASE PRECURSOR calcoaceticus]	5.2		
4476	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4477	X95455	G.gallus mRNA for RING zinc finger	9e-031	1321818	(X95455) RING zinc finger protein protein [Gallus	9e-038		

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					gallus]		
4478	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.13	<none></none>	<none></none>	<none></none>	
4479	J03607	Human 40-kDa keratin intermediate filament precursor gene.	0	1070608	keratin 19, type I, cytoskeletal - human sapiens]	9e-068	
4480	M90104	Human splicing factor SC35 mRNA, complete cds.	e-120	3929382	SPLICING FACTOR, ARGININE/SERI NE-RICH 10 (PUTATIVE MYELIN REGULATORY FACTOR 1) (MRF-1) >gi 555924 (U14648) putative myelin regulatory factor 1; MRF-1 [Mus musculus]	1.1	
4481	AF020762	Homo sapiens clone 1400 unknown protein mRNA, partial cds	6e-067	<none></none>	<none></none>	<none></none>	
4482	AE001386	Plasmodium falciparum chromosome 2, section 23 of 73 of the complete sequence	0.72	<none></none>	<none></none>	<none></none>	
4483		Pseudomonas aeruginosa autoinducer synthetase chloramphenicol- sensitive protein (rarD), and hypothetical protein (yafL) gene	0.005	1709793	SALIVARY PROLINE-RICH PROTEIN PO sapiens]	0.13	
4484	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

	<u> </u>	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4485	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4486	AE001406	Plasmodium falciparum chromosome 2, section 43 of 73 of the complete sequence	0.001	<none></none>	<none></none>	<none></none>	
4487	AE001417	Plasmodium falciparum chromosome 2, section 54 of 73 of the complete sequence	2.1	<none></none>	<none></none>	<none></none>	
4488	X90446	Canine herpesvirus DNA for ORF 1 (HSV1 UL44, EHV1 ORF 15 homolog) ORF2 (EHV1 ORF 16 homolog)	4.4	<none></none>	<none></none>	<none></none>	
4489	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.17	4008355	(Z68297) Similarity to Yeast TAT-binding homolog 7 (SW:TBP7_YEAS T); cDNA EST EMBL:D37124 comes from this gene; cDNA EST EMBL:D35150 comes from this gene; cDNA EST EMBL:D35400 comes from this gene; cDNA EST EMBL:D35400 comes from this gene; cDNA EST EMBL:D34900 comes >gi 4008373 gnl PI D e135984	3e-007	
4490	D78130	Homo sapiens mRNA for squalene epoxidase, complete cds	0	2443316	(D78130) squalene epoxidase [Homo sapiens]	5e-008	
4491	L18931	Buchnera aphidicola Arginyl tRNA synthetase	0.16	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		promoter region.					
		promoter region.					
4492	X17206	Human mRNA	e-112	1350976	40S	2e-005	
		for LLRep3			RIBOSOMAL		
					PROTEIN S2  >gi 939718		
4493	D28473	Human T-	e-157	440799	(U04953)	3e-005	
		lymphocyte		. 10723	isoleucyl-tRNA	30-003	
		mRNA for			synthetase [Homo		
		isoleucyl-tRNA			sapiens]		
		synthetase,					
4404	L13624	complete cds	2.6	AIONE	NOVE	110115	
4494	L13624	Cercopithecus aethiops C4	3.6	<none></none>	<none></none>	<none></none>	
		complement					
4495	M13011	Rat c-ras-H-1	0.25	<none></none>	<none></none>	<none></none>	
		gene, complete					
		cds.					
4496	Y10252	L.japonicus panC	0.38	627071	histidine-rich	4.4	
		gene			protein - Plasmodium		
1 1					lophurae		
4497	X76683	Plasmid vector	1e-093	987050	(X65335) lacZ	3e-015	
		рНМ2			gene product		
		betalactamase			[unidentified		
4400	3404406	gene		100065	cloning vector]		
4498	M24486	Human prolyl 4- hydroxylase	0	129365	PROLYL 4- HYDROXYLASE	2e-057	
		alpha subunit			ALPHA		
		mRNA, complete		i	SUBUNIT		
		cds, clone PA-11.			1.14.11.2) alpha		
					chain - chicken		
4499	D80004	Human mRNA for KIAA0182	2e-068	<none></none>	<none></none>	<none></none>	
		gene, partial cds					
4500	U22233	Human	0	<none></none>	<none></none>	<none></none>	
	I	methylthioadenos	Ĭ,	THOME	THOME	NONE	
		ine phosphorylase				1	
		(MTAP) mRNA,				[	
4501	DC2875	complete cds.		061440	(D (2055)		
4501	i	Human mRNA for KIAA0155	0	961442	(D63875)	2e-019	
		gene, complete			KIAA0155 gene product is related		
		cds > ::	ŀ		to C.elegans		
		gb G28541 G285			B0464.2 protein.		
	1	41 human STS			[Homo sapiens]	1	
		SHGC-31621.	502				

4.5	Nearest Neighbor			Nearest Neighbor			
1.5	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4502	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4503	<none> X85018</none>	<none> H.sapiens mRNA for UDP- GalNAc:polypept ide N- acetylgalactosami nyltransferase (T1)</none>	e-110	<none> 1709559</none>	POLYPEPTIDE N- ACETYLGALAC TOSAMINYLTR ANSFERASE (PROTEIN-UDP ACETYLGALAC TOSAMINYLTR ANSFERASE) N- ACETYLGALAC TOSAMINYLTR ANSFERASE) (GALNAC-T1) polypeptide N- acetylgalactosamin	2e-018	
					yltransferase [Rattus norvegicus]		
4504	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4505	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4506	AF067782	Papio hamadryas BC200 alpha scRNA gene, complete sequence	0.48	<none></none>	<none></none>	<none></none>	
4507	AF073298	Homo sapiens 4F5rel mRNA, complete cds	e-166	3641536	(AF073297) 4F5rel [Mus musculus] >gi 3641538 (AF073298) 4F5rel [Homo sapiens]	3e-013	
4508	M12922	Yeast (S.cerevisiae) chromosome III L terminal region DNA.	2e-010	188864	(M74027) mucin [Homo sapiens]	6e-023	
4509	X69524	M.squamata cabc1 mRNA for chlorophyll a/b/c binding protein precursor	1.3	<none></none>	<none></none>	<none></none>	
4510	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	1.2	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4512	U12404	Human Csa-19 mRNA, complete cds.	0	1709973	60S RIBOSOMAL PROTEIN L10A (CSA-19)	4e-056	
4513	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-014	<none></none>	<none></none>	<none></none>	
4514	<none></none>	<none></none>	<none></none>	121627	GLYCINE-RICH CELL WALL STRUCTURAL PROTEIN 1 PRECURSOR >gi 82244 pir  A26 099 glycine-rich cell wall structural protein - garden petunia >gi 20553 hybrida] >gi 225181 prf  12 10313A Gly rich structural protein [Petunia sp.]	2e-030	
4515	D87255	Hepatitis G virus RNA for polyprotein, complete cds	0.19	930045	(X15332) alpha-1 (III) collagen [Homo sapiens]	0.002	
4516	U31820	Gallus gallus Mel-1a melatonin receptor mRNA, complete cds.	3.3	1718187	ENVELOPE GLYCOPROTEIN GP340 glycoprotein 350/220 - human herpesvirus 4 >gi 59164 virus] >gi 306293 (L07923) glycoprotein 340	0.096	
4517	X68107	M.sativa msCHSII mRNA for chalcone synthase	3.4	<none></none>	<none></none>	<none></none>	
4518	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4519	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	6e-006	1065484	(U40415) similar to S. cerevisiae LAG1 (SP:P38703)	0.001	

	Nearest Neighbor			Nearest Neighbor			
*	•	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4520	D87671	Rat mRNA for TIP120, complete cds	1e-043	1799570	[D87671) TIP120 [Rattus norvegicus]	0.01	
4521	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4522	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4523	X16869	Human mRNA for elongation factor 1-alpha (clone CEF4)	4e-022	1085204	translation elongation factor eEF-1 alpha chain - zebra fish >gi 408805 (L23807) elongation factor 1-alpha [Danio rerio] >gi 454915 (X77689) translational elongation factor-1 alpha [Danio rerio] >gi 1009241 rerio] >gi 1091578 prf  2 021264A elongation fact	5.1	
4524	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-010	<none></none>	<none></none>	<none></none>	
4525	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>	
4526	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4527	AF069250	Homo sapiens okadaic acid- inducible phosphoprotein (OA48-18) mRNA, complete cds	7e-080	3037018	(AF041330) NADH dehydrogenase subunit 5 [Bodo saltans]	0.0001	
4528	AF069250	Homo sapiens okadaic acid- inducible phosphoprotein (OA48-18) mRNA, complete cds	7e-080	3037018	(AF041330) NADH dehydrogenase subunit 5 [Bodo saltans]	0.0001	

197	<u> </u>	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4529	U66532	Human beta4- integrin (ITGB4) gene, exons 7,8,9,10,11 and 12	0.51	119110	EBNA-1 NUCLEAR PROTEIN herpesvirus 4 (strain B95-8) >gi 1334880 (V01555) BKRF1 encodes EBNA-1 protein, latent cycle gene. [Human herpesvirus 4]	1e-023	
4530	X65319	Cloning vector pCAT-Enhancer	1e-074	987050	(X65335) lacZ gene product [unidentified cloning vector]	8e-011	
4531	AJ010841	Homo sapiens mRNA for putative thioredoxin-like protein	8e-028	3646128	(AJ010841) thioredoxin-like protein	0.062	
4532	D14034	Human gene for Zn-alpha2-glycoprotein, complete cds	0.005	<none></none>	<none></none>	<none></none>	
4533	M12670	Human fibroblast collagenase inhibitor mRNA, complete cds.	6e-098	1351250	METALLOPROT EINASE INHIBITOR.1 PRECURSOR (TIMP-1) >gi 1363927 pir  J C4303 matrix metalloproteinase-1 tissue inhibitor -baboon >gi 561546 hamadryas cynocephalus]	7e-008	
4534	M17196	A.californica (marine gastropod mollusc) neuropeptide gene (ganglion R14), exon 1, 5' end.	0.019	2135765	mucin 2 precursor, intestinal - human	0.003	
4535	AJ001454	Homo sapiens mRNA for testican-3	1.4	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
	`	astN vs. Genbank)		`		,		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4536	X75757	G.gallus cycB3 mRNA.	9e-040	729112	G2/MITOTIC- SPECIFIC CYCLIN B3	9e-019		
4537	Z27116	S.cerevisiae HBS1, MRP-L20 and PRP-16 genes	0.058	<none></none>	<none></none>	<none></none>		
4538	AF083322	Homo sapiens centriole associated protein CEP110 mRNA, complete cds	9e-051	1079393	chromokinesin - chicken >gi 603761 (U18309) chromokinesin [Gallus gallus]	0.012		
4539	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4540	M26325	Human cytokeratin 18 mRNA, 3' end.	0	125083	KERATIN, TYPE I CYTOSKELETA L 18 keratin 18, type I, cytoskeletal - human >gi 34037	2e-093		
4541	U37066	Human endogenous retrovirus strain XA38 pol polyprotein (pol) gene, partial cds	1.3	252486	P-selectin, CD62 [mice, Peptide, 768 aa] musculus]	1.8		
4542	Z30543	Turkey herpesvirus (HVT-delUs- Beta1 PKI3) gene for protein kinase	2e-027	<none></none>	<none></none>	<none></none>		
4543	M90077	Wheat translation elongation factor 1 alpha-subunit (TEF1) mRNA, complete cds.	0.14	<none></none>	<none></none>	<none></none>		
4544	AJ001235	Papio hamadryas ERV-9 like LTR insertion	2e-044	<none></none>	<none></none>	<none></none>		
4545	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4546	AF100654	Caenorhabditis elegans cosmid C24E9	0.41	<none></none>	<none></none>	<none></none>		
4547	L28821	Homo sapiens alpha mannosidase II isozyme mRNA, complete cds.	0	1679607	(X97650) myosin- I [Mus musculus]	4.5		

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4548	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-013	<none></none>	<none></none>	<none></none>	
4549	L20140	Zea mays pollen specific pectate lyase homologue gene, complete cds.	0.92	<none></none>	<none></none>	<none></none>	
4550	U33955	Human Down Syndrome region of chromosome 21, genomic sequence, clone A12H1-1F2.	4.4	<none></none>	<none></none>	<none></none>	
4551	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0005	<none></none>	<none></none>	<none></none>	
4552	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.042	<none></none>	<none></none>	<none></none>	
4553	X12660	Human chromosome 14 Ig JH (switch mu) DNA showing scattered homology to bcl2 gene exon 2 3'UTR	1e-006	2117245	(Z95586) hypothetical protein Rv1592c	2.1	
4554	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	284314	modulator recognition factor 1 - human factor I [Homo sapiens]	7.1	
4555	AF070523	Homo sapiens JWA protein mRNA, complete cds	0	3322740	(AE001222) conserved hypothetical protein [Treponema pallidum]	5.9	
4556	Z11900	H.sapiens OTF3 gene	0.13	<none></none>	<none></none>	<none></none>	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
4557	M24972	D.discoideum	4e-007	2605798	(AF027735) minor	5.30E-01	
		CT-rich satellite			ampullate silk		
		rDNA, clone			protein MiSp1		
		рСТ8.			[Nephila clavipes]		
4558	U95098	Xenopus laevis	8e-007	<none></none>	<none></none>	<none></none>	
1		mitotic					
		phosphoprotein			,		
		44 mRNA, partial					
1550	D22056	cds	0.06	<none></none>	<none></none>	<none></none>	
4559	D32056	Human gene for	0.00	\NONE>	\I\OI\I_>	THORIE	
		2-oxoglutarate dehydrogenase,					
		exon 1 sequence					
4560	AF034085	Caenorhabditis	0.025	1652167	(D90903)	4.8	
4360	AF034063	elegans UNC-45	0.023	1032107	hypothetical		
		(unc-45) gene,			protein		
		complete cds			Process		
4561	AF091242	Homo sapiens	0.0003	<none></none>	<none></none>	<none></none>	
4501	A1 0712-12	ATP	0.000				
		sulfurylase/APS					
		kinase 2 mRNA,					
		complete cds					
4562	M31520	Human ribosomal	1e-031	2072296	(U95098) mitotic	5.7	
		protein S24			phosphoprotein 44		
		mRNA.			[Xenopus laevis]		
4563	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4564	U95094	Xenopus laevis	0.0005	<none></none>	<none></none>	<none></none>	
		XL-INCENP			-		
		(XL-INCENP)					
		mRNA, complete					
45.55	A D 01 5 422	cds	4e-022	1665759	(D87432) Similar	5e-024	
4565	AB015432	Rattus norvegicus mRNA for LAT1	46-022	1003/39	to Schistosoma	JC-024	
		(L-type amino			mansoni amino		
		acid transporter			acid permease		
		1), complete cds			(L25068). [Homo		
		1), complete eds			sapiens]		
4566	AE001397	Plasmodium	0.0005	3875266	(Z77655)	5.90E+00	
.500	1	falciparum			predicted using		
		chromosome 2,	1		Genefinder;		
		section 34 of 73			similar to 7tm		
		of the complete			receptor		
		sequence			[Caenorhabditis		
					elegans]		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)  ACCESSION   DESCRIPTION   P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION		
4567	AE001397	Plasmodium falciparum chromosome 2, section 34 of 73 of the complete sequence	0.0005	3875266	(Z77655) predicted using Genefinder; similar to 7tm receptor [Caenorhabditis elegans]	5.90E+00	
4568	¥15155	Homo sapiens PHKB gene, exon 8, and repetitive elements	4e-033	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.7	
4569	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-03	2622750	(AE000921) DNA topoisomerase I [Methanobacteriu m thermoautotrophic um]	2.6	
4570	AE000688	Aquifex aeolicus section 20 of 109 of the complete genome	4.5	<none></none>	<none></none>	<none></none>	
4571	Z95123	Caenorhabditis elegans cosmid VZK8221, complete sequence [Caenorhabditis elegans]	0.4	<none></none>	<none></none>	<none></none>	
4572	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3.00E-08	<none></none>	<none></none>	<none></none>	
4573	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.3	
4574	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-006	<none></none>	<none></none>	<none></none>	
4575	U18671	Human Stat2 gene, complete cds.	2e-023	728831	IIII ALU SUBFAMILY J WARNING ENTRY	0.002	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
4576	Z83241	Caenorhabditis elegans cosmid T25C8, complete sequence [Caenorhabditis elegans]	1.1	1176988	IOLD PROTEIN protein [Bacillus subtilis] >gi 2636519 gnl PI D e1184698 catabolism [Bacillus subtilis]	5.3	
4577	L04690	Cricetulus griseus cholesterol 7- alpha- hydroxylase gene, complete cds. > :: gb I26617 I26617 Sequence 35 from patent US 5558999 > :: gb AR008072 AR 008072 Sequence 35 from patent US 5753431	3.2	212906	(L02621) intestinal zipper protein [Gallus gallus]	4.1	
4578	Z54191	A.pleuropneumon iae tfbB gene encoding transferrin receptor.		2102696	(U72761) karyopherin beta 3 [Homo sapiens]	8.6	
4579	X17025	Human homolog of yeast IPP isomerase > :: gb G27043 G270 43 human STS SHGC-31614.	2e-035	<none></none>	<none></none>	<none></none>	
4580	L32977	Homo sapiens (clone f17252) ubiquinol cytochrome c reductase Rieske iron-sulphur protein (UQCRFS1) gene, exon 2	0.00E+00		UBIQUINOL- CYTOCHROME C REDUCTASE IRON-SULFUR SUBUNIT PRECURSOR (RIESKE IRON- SULFUR PROTEIN) (RISP) >gi 488299 (L32977) Rieske Fe-S protein		
4581	M26708	Human prothymosin alpha mRNA (ProT-alpha), complete cds.	0	190369	(J04798) open reading frame A; putative [Homo sapiens]	6e-018	

	7	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4582	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-014	2314130	(AE000607) H. pylori predicted coding region HP0985	3.3		
4583	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	1236083	(U49507) Lisch7 [Mus musculus]	4.3		
4584	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-014	348196	(L19917) immunoglobulin heavy-chain subgroup VIII V- D-J region [Homo sapiens]	9.7		
4585	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4586	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4587	X52601	H.sapiens hTOP1 gene for topoisomerase, 5'end	4.6	<none></none>	<none></none>	<none></none>		
4588	AF038604	Caenorhabditis elegans cosmid B0546	0.17	<none></none>	<none></none>	<none></none>		
4589	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4590	U23 <u>441</u>	Tetrahymena thermophila B internal deletion sequence.	0.0005	1469281	(U08801) envelope glycoprotein [Human immunodeficiency virus type 1]	1.1		
4591	AC005276	Homo sapiens clone fragment UWGC:gap3 from 7q31.3, complete sequence [Homo sapiens]	0.009	<none></none>	<none></none>	<none></none>		
4592	D84117	Homo sapiens DNA for prostacyclin synthase, exon 3	0.48	<none></none>	<none></none>	<none></none>		
4593	U28153	Caenorhabditis elegans UNC-76 (unc-76) gene, complete cds.	1.30E-01	<none></none>	<none></none>	<none></none>		

	Nearest Neighbor			Nearest Neighbor			
	,	astN vs. Genbank)			. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4594	U67274	Human metastasis suppressor (KAI1) gene, exon 1, and complete cds	1e-008	<none></none>	<none></none>	<none></none>	
4595	AF009621	Onchocerca volvulus cytosolic Cu/Zn superoxide dismutase (OvSOD1) and extracellular Cu/Zn superoxide dismutase (OvSOD2) genes, complete cds	4	<none></none>	<none></none>	<none></none>	
4596	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4597	<none></none>	<none></none>	<none></none>	2078483	(U43200) antifreeze glycopeptide AFGP polyprotein precursor [Boreogadus saida]	0.78	
4598	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4599	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4600	AL021806	Homo sapiens DNA sequence from PAC 779B17 on chromosome 22q13.1. Contains exon trap, complete sequence	4e-029	728836	!!!! ALU - SUBFAMILY SP WARNING ENTRY	0.002	
4601	AL022222	Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS *** from contig 3-118, complete sequence	4.9	<none></none>	<none></none>	<none></none>	
4602	Z73149	N.tabacum DNA (recombination breakpoint between T-DNA and plant DNA)	1.6	<none></none>	<none></none>	<none></none>	

	<u> </u>	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
4603	AF082835	Mus spretus E6-	4	<none></none>	<none></none>	<none></none>	
		AP ubiquitin-					
		protein ligase	2 000	500000	1111 47 77		
4604	AF050123	Homo sapiens	3e-009	728838	!!!! ALU SUBFAMILY SX	6.7	
		hypoxia-inducible factor 1 alpha			WARNING		
		subunit (HIF1A)			ENTRY		
		gene, exon 10					
4605	U95102	Xenopus laevis	7e-006	<none></none>	<none></none>	<none></none>	
		mitotic					
		phosphoprotein					
		90 mRNA,					
		complete cds	0.1	2041726	TRANSCRIPTIO	9.0	
4606	AF001355	Pseudomonas	2.1	3041736	TRANSCRIPTIO N FACTOR SOX-	8.9	
		syringae pv. syringae DNA			11		
		binding protein		ł	^ ^		
		HpkR (hpkR),					
		histidine protein					
		kinase HpkY					
		(hpkY),					
		phosphate					
		acceptor regulatory protein					
		CheY-2 (cheY-2),					
		ankyrin AnkF		İ			
:		(ankF), and					
		catalase isozyme		]	•		
		catalytic subuni	0.005.00	2122155	INVENTIONAL	0.007	
4607	U95102	Xenopus laevis mitotic	8.00E-08	3123155	HYPOTHETICAL 49.0 KD TRP-	2e-027	
		phosphoprotein			ASP REPEATS		
		90 mRNA.			CONTAINING		
		complete cds			PROTEIN		
					F55F8.5 IN		
					CHROMOSOME		
			•		I family [Caenorhabditis		
					elegans]		
4608	<none></none>	<none></none>	<none></none>	1170978	MYOCYTE	0.18	
17000	110112	-1101112	1,01,0	********	NUCLEAR		
					FACTOR (MNF)		
					musculus]		
4609	U95098	Xenopus laevis	4e-009	2072296	(U95098) mitotic	8.9	
		mitotic			phosphoprotein 44		
		phosphoprotein 44 mRNA, partial			[Xenopus laevis]		
		cds		ŀ			
L	<u> </u>	1000	<u> </u>	<u> </u>	l		

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4610	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>	
4611	X75861	H.sapiens TEGT gene	e-177	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.8	
4612	U19867	Cloning vector pSPL3, exon splicing vector, complete sequence, HIV envelope protein gp160 and beta- lactamase, complete cds.	5e-055	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-011	
4613	U73332	Human non- coding genomic sequence upstream from unique L0 sequence in the alpha-globin gene cluster	8e-008	<none></none>	<none></none>	<none></none>	
4614	<none></none>	<none></none>	<none></none>	193952	(J03770) homeobox protein [Mus musculus]	6	
4615	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	586875	HYPOTHETICAL 29.2 KD PROTEIN IN METS-KSGA INTERGENIC REGION >gi 2127033 pir  S 66068 hypothetical protein - Bacillus subtilis subtilis] >gi 2632306 gnl PI D e1181972 (Z99104) similar to hypothetical proteins [Bacillus subtilis]		
4616	K00384	Yeast (S.cerevisiae) mitochondrial var1 gene, 5'	0.001	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		flank.					
4617	J04628	Rattus norvegicus 3-hydroxyiso- butyrate mRNA, 3' end.	e-154	416873	3- HYDROXYISOB UTYRATE DEHYDROGENA SE PRECURSOR (HIBADH) >gi 111295 pir  A3 2867 3- hydroxyisobutyrat e dehydrogenase (EC 1.131) precursor - rat (fragment) >gi 556389 (J04628) 3- hydroxyisobutyrat e dehydrogenase [Rattus norvegicus] <none></none>	1e-049	
		mitotic phosphoprotein 44 mRNA, partial cds		NONE	- NOINE	<nune></nune>	
4619	U10361	Saccharomyces cerevisiae Snf8p (SNF8) gene, complete cds.	2.7	<none></none>	<none></none>	<none></none>	
4620	D42044	Human mRNA for KIAA0090 gene, partial cds	e-151	577301	(D42044) The ha3523 gene product is related to S.cerevisiae gene product located in chromosome III. [Homo sapiens]	4e-052	
4621		Saccharomyces cerevisiae Snf8p (SNF8) gene, complete cds.	2.7	<none></none>	<none></none>	<none></none>	
4622	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

	N	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
4623	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3.00E-10	<none></none>	<none></none>	<none></none>	
4624	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3.00E-10	<none></none>	<none></none>	<none></none>	
4625	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4626	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4627	X06747	Human hnRNP core protein A1	7e-049	87650	heterogeneous ribonuclear particle protein A1.beta - human >gi 36102 (X06747) protein A1-alpha (AA 1- 320) [Homo sapiens]	6e-005	
4628	X03559	Human mRNA for F1-ATPase beta subunit (F-1 beta) > :: dbj D00022 HUM F1B Homo sapiens mRNA for F1 beta subunit, complete cds	e-100	114549	ATP SYNTHASE BETA CHAIN, MITOCHONDRI AL PRECURSOR >gi 106207 pir  A3 3370 H+- transporting ATP synthase (EC 3.6.1.34) beta chain precursor, mitochondrial - human >gi 179281 (M27132) ATP synthase beta subunit precursor [Homo sapiens]	2e-024	
4629	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4630	K00915	paramecium species 1,168 mt dna dimer: replication init. region.	7.00E-05	<none></none>	<none></none>	<none></none>	
4631	K00915	paramecium species 1,168 mt dna dimer: replication init. region.	7.00E-05	<none></none>	<none></none>	<none></none>	
4632	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

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SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4633	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4634	Z28261	S.cerevisiae chromosome XI reading frame ORF YKR036c	0.042	417748	PROTEIN TRANSPORT PROTEIN SEC13	0.0002	
4635	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
4636	AF088034	Homo sapiens full length insert cDNA clone ZC24F03	0	854598	(X87611) ORF YJR83.18 [Saccharomyces cerevisiae]	2e-024	
4637	M83094	Homo sapiens cytosolic selenium-dependent glutathione peroxidase gene, complete cds, and rhoh12 gene, 3' end.	3.00E-08	<none></none>	<none></none>	<none></none>	
4638	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, .complete cds	2e-006	1176711	HYPOTHETICAL 21.6 KD PROTEIN F37A4.2 IN CHROMOSOME III >gi 1078851 pir  S 44639 F37A4.2 protein - Caenorhabditis elegans >gi 458960	2e-017	
4639	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	1176711	HYPOTHETICAL 21.6 KD PROTEIN F37A4.2 IN CHROMOSOME III >gi 1078851 pir  S 44639 F37A4.2 protein - Caenorhabditis elegans >gi 458960	2e-017	

	Nearest Neighbor			Nearest Neighbor			
ana		lastN vs. Genbank)	T	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4640	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
4641	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
4642	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	4056582	(AF039530) RepA [Egyptian sugarcane streak virus]	3.4	
4643	U96174	Onchocerca volvulus OvB8 mRNA, partial cds	3.2	<none></none>	<none></none>	<none></none>	
4644	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4645	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-005	3236220	(U62541) immunoreactive 14 kDa protein BA14k [Brucella abortus]	4.5	
4646	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-005	3236220	(U62541) immunoreactive 14 kDa protein BA14k [Brucella abortus]	4.5	
4647	AL010224	Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS *** from contig 4-04, complete sequence	0.003	2492906	ANNEXIN VII (SYNEXIN) frog >gi 790544 (U16365) annexin VII [Xenopus laevis]	1.4	
4648	L39413	Atractylodes japonica chloroplast NADH dehydrogenase (ndhF) gene, complete cds	0.003	<none></none>	<none></none>	<none></none>	
4649	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete	4e-013	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor			Nearest Neighbor			
	, , , ,	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
	•	cds						
4650	U79403	Meleagris gallopavo microsatellite repeat sequence	0.46	2498691	OUTER DENSE FIBER PROTEIN bovine >gi 1165006 (X69514) outer dense fiber protein protein [Bos taurus]	1.4		
4651	U27780	Stealth virus 1 clone C16138 T3.1	2	<none></none>	<none></none>	<none></none>		
4652	U27780	Stealth virus 1 clone C16138 T3.1	2	<none></none>	<none></none>	<none></none>		
4653	U78817	Saccharomyces cerevisiae killer virus M1, complete genome	0.026	<none></none>	<none></none>	<none></none>		
4654	U78817	Saccharomyces cerevisiae killer virus M1, complete genome	0.026	<none></none>	<none></none>	<none></none>		
4655	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
4656	X07036	Human mRNA stimulatory GTP- binding protein alpha subunit	3e-071	232142	GUANINE NUCLEOTIDE- BINDING PROTEIN G(S), ALPHA SUBUNIT (ADENYLATE CYCLASE- STIMULATING G ALPHA PROTEIN) >gi 71886 pir  RG PGA2 GTP- binding regulatory protein Gs alpha-2 chain (adenylate cyclase- stimulating) - pig >gi 1958 (X63893) alpha-stimulatory subunit	8e-027		

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4657	L05586	Kinetoplast Trypanosoma brucei (IsTaR 1 serodeme) putative NADH dehydrogenase subunit (nd9) mRNA, complete cds.	0.0001	4063042	(AF068065) GP900; mucin-like glycoprotein [Cryptosporidium parvum]	0.19	
4658	AF044763	Cecropis ariel microsatellite HrU6 allele 1 repeat region	3e-006	<none></none>	<none></none>	<none></none>	
4659	X82630	A.longa plastid rps12, orf126 and orf288 genes	0.22	<none></none>	<none></none>	<none></none>	
4660	U68098	Human poly(A)- binding protein (PABP) gene, exons 6 and 7	0.023	<none></none>	<none></none>	<none></none>	
4661	U68098	Human poly(A)- binding protein (PABP) gene, exons 6 and 7	0.023	<none></none>	<none></none>	<none></none>	
4662	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	1022683	[Rattus norvegicus]	1.4	
4663	M15353	Homo sapiens cap-binding protein mRNA, complete cds	0	<none></none>	<none></none>	<none></none>	
4664	Z57610	H.sapiens CpG DNA, clone 187a10, reverse read cpg187a10.rt1a.	3e-048	417134	HEPATOCYTE NUCLEAR FACTOR 3-BETA norvegicus]	2.00E-10	
4665	L11707	Hevea brasiliensis Mn- superoxide dismutase (SODMn) gene, complete cds.	2.6	<none></none>	<none></none>	<none></none>	
4666	D42073	Human mRNA for reticulocalbin, complete cds	3e-019	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.4	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4667	L12350	Human thrombospondin 2 (THBS2) mRNA, complete cds.	0	<none></none>	<none></none>	<none></none>	
4668	L11707	Hevea brasiliensis Mn- superoxide dismutase (SODMn) gene, complete cds.	2.6	<none></none>	<none></none>	<none></none>	
4669	AC000043	Homo sapiens Chromosome 22q13 Cosmid Clone p74a8, complete sequence [Homo sapiens]	2e-016	134589	TRANSCRIPTION REGULATORY PROTEIN SNF2 SWI2) (REGULATORY PROTEIN GAM1) (TRANSCRIPTION FACTOR TYE3) >gi 101629 pir  S1 5047 SNF2 protein - yeast protein [Saccharomyces cerevisiae] >gi 172632 (M61703) SNF2protein [Saccharomyces cerevisiae] cerevisiae] >gi 127	1.5	
4670	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	69700	interleukin-1 beta precursor - bovine	0.6	
4671	U44975	Homo sapiens DNA-binding protein CPBP (CPBP) mRNA, partial cds	2e-045	1848233	(U44975) DNA- binding protein CPBP [Homo sapiens]	0.009	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4672	AF038406	Homo sapiens NADH dehydrogenase- ubiquinone Fe-S protein 8 23 kDa subunit (NDUFS8) gene, nuclear gene encoding mitochondrial protein, complete cds	0	2326168	(U32107) type VII collagen [Mus musculus]	1.5	
4673	X67951	H.sapiens mRNA for proliferation- associated gene	0	548453	THIOREDOXIN PEROXIDASE 2 CELL ENHANCING FACTOR A) (NKEF-A) >gi 423025 pir  A4 6711 proliferation associated gene (pag) protein - human gene product [Homo sapiens]	2e-083	
4674	AC001013	Homo sapiens (subclone 2_d1 from P1 H43) DNA sequence	2e-017	2072961	(U93568) putative p150 [Homo sapiens]	0.0001	
4675	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	1589837	(U68729) cuticle preprocollagen [Meloidogyne incognita]	0.035	
4676	M15353	Homo sapiens cap-binding protein mRNA, complete cds	0	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4677	M37583	Human histone (H2A.Z) mRNA, complete cds.	0	121994	HISTONE H2A.Z  >gi 89608 pir  S03 642 histone  H2A.Z - bovine  >gi 92380 pir  S03 644 histone  H2A.Z - rat  >gi 106267 pir  A3 5881 histone  H2A.Z - human  sapiens] >gi 57808  (X52316) histone  H2A.Z (AA 1-  127) taurus]  >gi 184060  (M37583) histone  (H2A.Z) [Homo  sapien	1e-055	
4678	M15353	Homo sapiens cap-binding protein mRNA, complete cds	0	<none></none>	<none></none>	<none></none>	
4679	Z57610	H.sapiens CpG DNA, clone 187a10, reverse read cpg187a10.rt1a.	4e-094	404764	(L10409) fork head related protein [Mus musculus]	4e-024	
4680	Z57610	H.sapiens CpG DNA, clone 187a10, reverse read cpg187a10.rt1a.	4e-094	404764	(L10409) fork head related protein [Mus musculus]	4e-024	
4681	Z57610	H.sapiens CpG DNA, clone 187a10, reverse read cpg187a10.rt1a.	4e-094	404764	(L10409) fork head related protein [Mus musculus]	4e-024	
4682	L11707	Hevea brasiliensis Mn- superoxide dismutase (SODMn) gene, complete cds.	2.6	<none></none>	<none></none>	<none></none>	
4683	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4684	<none></none>	<none></none>	<none></none>	2114323	(D88734) membrane glycoprotein [Equine herpesvirus 1]	0.052	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4685	AJ224875	Homo sapiens mRNA for putative glucosyltransferas e, partial cds	0	2996578	(AJ224875) glucosyltransferas e [Homo sapiens]	e-118	
4686	AB019534	Homo sapiens gene for cathepsin L2, complete cds	2e-045	<none></none>	<none></none>	<none></none>	
4687	J03799	Human colin carcinoma laminin-binding protein mRNA, complete cds.	e-166	34272	(X15005) pot. lamimin-binding protein (AA 1 - 300) [Homo sapiens]	5e-032	
4688	<none></none>	<none></none>	<none></none>	2114323	(D88734) membrane glycoprotein [Equine herpesvirus 1]	0.052	
4689	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	9e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	9.8	
4690	D44598	Saccharomyces cerevisiae chromosome VI phage 4121	1e-009	3947877	(AL034382) putative mitosis and maintenance of ploidy protein [Schizosaccharom yces pombe]	6e-061	
4691	AF053520	Homo sapiens allele 12 fragile site locus	0.61	<none></none>	<none></none>	<none></none>	
4692	D16195	Mouse gene for acrogranin precursor, complete cds	0.059	<none></none>	<none></none>	<none></none>	
4693	U90904	Human clone 23773 mRNA sequence	0	3130153	(AB008857) calcium2+ sensing receptor	1.5	
4694	L22398	Homo sapiens DNA sequence, repeat region.	7e-017	987050	(X65335) lacZ gene product [unidentified cloning vector]	0.1	
4695	L22398	Homo sapiens DNA sequence, repeat region.	7e-017	987050	(X65335) lacZ gene product [unidentified cloning vector]	0.1	

	<u> </u>	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)	:	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4696	J03746	Human glutathione S- transferase mRNA, complete cds.	e-170	121740	GLUTATHIONE S- TRANSFERASE, MICROSOMAL >gi 87562 pir  B28 083 glutathione transferase glutathione S- transferase [Homo sapiens] >gi 1195483 sapiens] >gi 1621433 (U71213) microsomal glutathione s- transferase [Homo sapiens]	2e-038	
4697	AF082283	Homo sapiens CARD- containing apoptotic signaling protein (BCL10) mRNA, complete cds	5e-046	4049460	(AJ006288) bcl-10 [Homo sapiens] signaling protein [Homo sapiens]	0.005	
4698	D64142	Human mRNA for histone H1x, complete cds	1e-039	<none></none>	<none></none>	<none></none>	
4699	AB001899	Homo sapiens PACE4 gene, exon 2	4e-012	3860844	(AJ235271) NADH DEHYDROGENA SE I CHAIN L	3.5	
4700	X16869	Human mRNA for elongation factor 1-alpha (clone CEF4)	0	1169475	ELONGATION FACTOR 1- ALPHA 1	6e-061	
4701	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6.00E-05	<none></none>	<none></none>	<none></none>	
4702	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-013	2501465	PROBABLE UBIQUITIN CARBOXYL- TERMINAL HYDROLASE FAM (UBIQUITIN THIOLESTERAS	0.0003	

	N	learest Neighbor	:	Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					E FAM)		
4703	D44598	Saccharomyces cerevisiae chromosome VI phage 4121	1e-009	3947877	(AL034382) putative mitosis and maintenance of ploidy protein [Schizosaccharom yces pombe]	6e-061	
4704	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	6e-006	<none></none>	<none></none>	<none></none>	
4705	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.4	
4706	AB001899	Homo sapiens PACE4 gene, exon 2	4e-012	3860844	(AJ235271) NADH DEHYDROGENA SE I CHAIN L	3.4	
4707	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4708	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-008	<none></none>	<none></none>	<none></none>	
4709	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	6.40E+00	
4710	L39064	Homo sapiens interleukin 9 receptor precursor (IL9R) gene, complete cds	1e-006	4063042	(AF068065) GP900; mucin-like glycoprotein	1e-006	
4711	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.0002	331908	(K02714) envelope polyprotein [Friend murine leukemia virus]	8	

		Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)  ACCESSION   DESCRIPTION   P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION			
4712	AF065249	Entodinium caudatum 14-3-3 protein mRNA, partial cds	1	<none></none>	<none></none>	<none></none>	
4713	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	7.9	
4714	<none></none>	<none></none>	<none></none>	186396	(M94131) mucin [Homo sapiens]	2.5	
4715	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-009	<none></none>	<none></none>	<none></none>	
4716	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4717	Z56314	H.sapiens CpG DNA, clone 10h10, reverse read cpg10h10.rt1a.	4e-012	2444024	(U77782) N- methyl-D- aspartate receptor 2C subunit precursor [Homo sapiens]	9.8	
4718	D55696	Human mRNA for cysteine protease, complete cds	e-113	2842759	LEGUMAIN PRECURSOR (ASPARAGINYL ENDOPEPTIDAS E) >gi 1743266 gnl PI D e286211 (Y09862) legumain [Homo sapiens]	1e-006	
4719	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	9e-008	<none></none>	NONE>	<none></none>	
4720	D63480	Human mRNA for KIAA0146 gene, partial cds	0	1469874	(D63480) The KIAA0146 gene product is novel. [Homo sapiens]	2e-079	
4721	AB001579	Rice dwarf virus genomic RNA, segment 2, complete sequence	1.3	<none></none>	<none></none>	<none></none>	
4722	<none></none>	<none></none>	<none></none>	3873550	(AL033534) serine-rich protein	2.7	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4723	AL010156	Plasmodium falciparum DNA *** SEQUENCING	0.77	<none></none>	<none></none>	<none></none>	
		IN PROGRESS  *** from contig  3-87, complete sequence					
4724	AF059198	Homo sapiens protein kinase/endoribon ulcease	2	119110	EBNA-1 NUCLEAR PROTEIN herpesvirus 4 (strain B95-8) >gi 1334880 (V01555) BKRF1 encodes EBNA-1 protein, latent cycle gene. [Human herpesvirus 4]	8e-007	
4725	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4727	D38616	Human mRNA for phosphorylase kinase alpha subunit, complete cds	3.5	3522948	(AC004411) hypothetical protein [Arabidopsis thaliana]	0.18	
4728	D38616	Human mRNA for phosphorylase kinase alpha subunit, complete cds	3.5	3522948	(AC004411) hypothetical protein [Arabidopsis thaliana]	0.18	
4729	Z11808	T.glis interphotorecepto r retinoid binding protein gene, exon 1	1.6	<none></none>	<none></none>	<none></none>	
4730	AF065988	Homo sapiens keratocan gene, complete cds	1.4	<none></none>	<none></none>	<none></none>	
4731	X60026	M.domesticus small nuclear 4.5 S RNA gene	0.0003	2853301	(AF007194) mucin [Homo sapiens]	5.5	
4732	M13793	Mouse 56 kdal protein mRNA from an interferon activated gene, exon 1, 5' end.	0.3	136814	HYPOTHETICAL PROTEIN UL11 RL11 FAMILY [Human cytomegalovirus]	2.3	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID			110	2042750	LECUMARI	1e-006	
4733	D55696	Human mRNA	e-113	2842759	LEGUMAIN PRECURSOR	16-000	
		for cysteine			(ASPARAGINYL		
		protease, complete cds			ENDOPEPTIDAS		
		complete cus			E)		
					>gi 1743266 gnl PI		
					D e286211		
					(Y09862)		
					legumain [Homo		
					sapiens]	270275	
4734	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4735	<none></none>	<none></none>	<none></none>	322647	glycine-rich protein GRP22 -	3e-021	
					protein GRP22 -   rape > gi   17821		
1726	AIONES	<none></none>	<none></none>	188864	(M74027) mucin	0.002	
4736	<none></none>	<none></none>	NONE	100004	[Homo sapiens]	0.002	
4737	U95102	Xenopus laevis	2e-005	<none></none>	<none></none>	<none></none>	
4/3/	093102	mitotic	20-003	110112			
1		phosphoprotein					
1		90 mRNA,					
		complete cds					
4738	AB018270	Homo sapiens	0	2072296	(U95098) mitotic	1.8	
		mRNA for			phosphoprotein 44		
		KIAA0727			[Xenopus laevis]		
		protein, partial					
4739	AB018270	Homo sapiens	0	2072296	(U95098) mitotic	1.8	
7/37	ABOTOZIO	mRNA for			phosphoprotein 44		
		KIAA0727			[Xenopus laevis]		
		protein, partial					
l		cds				110117	
4740	AE001382	Plasmodium	0.25	<none></none>	<none></none>	<none></none>	
		falciparum					
		chromosome 2, section 19 of 73					
		of the complete					
		sequence					
4741	AE001382	Plasmodium	0.25	<none></none>	<none></none>	<none></none>	
'''		falciparum					
		chromosome 2,					
		section 19 of 73					
		of the complete					
		sequence		<u> </u>			

	Nearest Neighbor			Nearest Neighbor			
	,	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4742	X55038	Mouse mCENP-B gene for centromere autoantigen B	0.001	3879362	(Z81113) similar to DnaJ, prokaryotic heat shock protein, Zinc finger, C2H2 type; cDNA EST yk290e12.5 comes from this gene; cDNA EST yk290e12.3 comes from this gene; cDNA EST yk447h4.5 comes from this gene; cDNA EST yk447h4.5 comes from this gene; cDNA EST yk474e4	7e-007	
4743	AF054024	Rattus norvegicus polymorphic marker D9UIA2 sequence	0.62	<none></none>	<none></none>	<none></none>	
4744	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0005	<none></none>	<none></none>	<none></none>	
4745	Z11808	T.glis interphotorecepto r retinoid binding protein gene, exon 1	1.6	<none></none>	<none></none>	<none></none>	
4746	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4747	AF047470	Homo sapiens malate dehydrogenase precursor complete cds	1e-019	2995307	(AL022268) putative aminotransferase	0.12	
4748	AF029890	Homo sapiens hepatitis B virus X interacting protein (XIP) mRNA, complete cds	e-161	2745883	(AF029890) hepatitis B virus X interacting protein [Homo sapiens]	2e-044	

,	N	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4750	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-008	1723019	HYPOTHETICAL 29.6 KD PROTEIN CY251.12C >gi 1405764 gnl PI D e249453 (Z74410) hypothetical protein Rv0093c [Mycobacterium tuberculosis]	2.5		
4751	M37583	Human histone (H2A.Z) mRNA, complete cds.	0	121994	HISTONE H2A.Z >gi 89608 pir  S03 642 histone H2A.Z - bovine >gi 92380 pir  S03 644 histone H2A.Z - rat >gi 106267 pir  A3 5881 histone H2A.Z - human sapiens] >gi 57808 (X52316) histone H2A.Z (AA 1- 127) taurus] >gi 184060 (M37583) histone (H2A.Z) [Homo sapien	1e-055		
4752	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-014	<none></none>	<none></none>	<none></none>		
4753	X65279	pWE15 cosmid vector DNA	7e-079	987050	(X65335) lacZ gene product [unidentified cloning vector]	1e-013		
4754	D38549	Human mRNA for KIAA0068 gene, partial cds	e-169	<none></none>	<none></none>	<none></none>		
4755	L27835	Pangasianodon gigas growth hormone (GH) mRNA, complete cds.	1.5	538251	(D00322) polyprotein [Tomato black ring virus]	5.8		

		Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		,	s. Non-Redundant Pr			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
4756	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	1477565	(U50078) p619 [Homo sapiens]	8.9		
4757	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	1477565	(U50078) p619 [Homo sapiens]	8.9		
4758	U47414	Human cyclin G2 mRNA, complete cds	e-116	<none></none>	<none></none>	<none></none>		
4759	AB014560	Homo sapiens mRNA for KIAA0660 protein, complete cds	e-173	<none></none>	<none></none>	<none></none>		
4760	L35664	Homo sapiens (subclone H8 8_f5 from P1 35 H5 C8) DNA sequence.	1e-030	2072966	(U93570) p40 [Homo sapiens]	0.001		
4761	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.1		
4762	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-013	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.1		
4763	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	<none></none>	<none></none>	<none></none>		
4764	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-012	<none></none>	<none></none>	<none></none>		
4765	M59317	Mouse low affinity IgE receptor (FceRII) gene sequence.	1e-006	2135765	mucin 2 precursor, intestinal - human	0.0003		

	1	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4766	D14034	Human gene for Zn-alpha2- glycoprotein, complete cds	3e-008	119379	RETROVIRUS- RELATED ENV POLYPROTEIN	6e-007	
4767	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4768	M61185	Bovine glutamic acid-rich protein mRNA, complete cds.	0.01	2781362	(AC003113) F24O1.18 [Arabidopsis thaliana]	1.1	
4769	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4770	Z62012	H.sapiens CpG DNA, clone 61g4, reverse read cpg61g4.rt1a	0.076	1582765	YFW1 gene [Saccharomyces cerevisiae]	2.9	
4771	M29065	Human hnRNP A2 protein mRNA.	0	4049652	(AF063866) ORF MSV017 hypothetical protein [Melanoplus sanguinipes entomopoxvirus]	5.9	
4772	D12525	Homo sapiens cytochrome P450IA1 gene, 3'flanking region	6e-016	728837	!!!! ALU SUBFAMILY SQ WARNING ENTRY	9.6	
4773	M16660	Human 90-kDa heat-shock protein gene, cDNA, complete cds.	e-109	2119731	HSP90 - mouse (fragment) protein {C-terminal} [mice, heart, Peptide Partial, 194 aa] [Mus sp.]	1e-023	
4774	AF043105	Homo sapiens glutathione S- transferase mu 3	9e-020	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.63	
4775	U43374	Human normal keratinocyte mRNA.	0	120179	FINQ PROTEIN >gi 73172 pir  BV ECFQ finQ protein - Escherichia coli plasmid R820a	9	
4776	U00684	Human unknown mRNA.	2e-014	2224667	(AB002361) KIAA0363 [Homo sapiens]	6.6	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4777	M22299	Human T-plastin polypeptide mRNA, complete cds, clone p4. > :: gb I08151  Sequence 1 from Patent EP 0345726	4e-008	<none></none>	<none></none>	<none></none>	
4778	M95623	Homo sapiens hydroxymethylbil ane synthase gene, complete cds.	3e-018	3002527	(AF010144) neuronal thread protein AD7c- NTP [Homo sapiens]	0.52	
4779	X52329	pBluescript II KS(-) vector DNA, phagemid excised from lambda ZAPII	0	2117615	catalase - Campylobacter jejuni	2e-009	
4780	X52329	pBluescript II KS(-) vector DNA, phagemid excised from lambda ZAPII	0	2117615	catalase - Campylobacter jejuni	2e-009	
4781	AF061034	Homo sapiens FIP2 alternatively translated mRNA, complete cds	0	3127084	(AF061034) FIP2 [Homo sapiens]	9e-089	
4782	Z64776	H.sapiens CpG DNA, clone 167d8, forward read cpg167d8.ft1b.	0.0002	1777782	(U52513) ISG family member [Homo sapiens]	1.8	
4783	D31786	Acyrthosiphon kondoi endosymbiont DNA, S10 and spc ribosomal protein gene operons, complete and partial cds	1.1	2134310	cell division control protein CDC37 homolog splice form 1 - chicken	4e-005	
4784	L05491	Homo sapiens T- plastin gene, last exon (16).	0	2506254	T-PLASTIN	3e-018	
4785	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	8e-007	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4786	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-006	3877438	(Z72510) similar to G-protein coupled receptor [Caenorhabditis elegans]	2	
4787	L38250	Mycoplasma penetrans p35 lipoprotein and p33 lipoprotein genes, complete cds	0.041	<none></none>	<none></none>	<none></none>	
4788	J03537	Human ribosomal protein S6 mRNA, complete cds.	e-138	133978	40S RIBOSOMAL PROTEIN S6 protein S6 - rat >gi 70933 pir  R3 MS6 ribosomal protein S6 - mouse >gi 319910 pir  R3 HU6 ribosomal protein S6 - human >gi 36148 (X67309) ribosomal protein S6 [Homo sapiens] >gi 54010 (Y00348) ribosomal protein S6 [Mus musculus] >g	3e-033	
4789	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	2.6	
4790	AF041210	Homo sapiens midline 1 fetal kidney isoform 3	0.41	<none></none>	<none></none>	<none></none>	
4791	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.2	

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		`	s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4792	S60885	LYAR=cell growth regulating nucleolar protein	2e-026	2498524	CELL GROWTH REGULATING NUCLEOLAR PROTEIN >gi 423488 pir  A4 0683 cell growth regulating nucleolar protein LYAR - mouse >gi 300372 bbs 13 1782	0.43	
4793	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4794	U28687	Human zinc finger containing protein ZNF157	3e-027	1731444	ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2) >gi 1020145 (M27878) DNA binding protein	3e-008	
4795	AF086438	Homo sapiens full length insert cDNA clone ZD80G11	0.0002	<none></none>	<none></none>	<none></none>	
4796	L28997	Homo sapiens ARL1 mRNA, complete cds	3e-006	<none></none>	<none></none>	<none></none>	
4797	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-008	1280126	(U55375) K03E6.4 [Caenorhabditis elegans]	2e-012	
4798	AE001415	Plasmodium falciparum chromosome 2, section 52 of 73 of the complete sequence	0.015	<none></none>	<none></none>	<none></none>	
4799	D21853	Human mRNA for KIAA0111 gene, complete cds	0	729821	EUKARYOTIC INITIATION FACTOR 4A- LIKE NUK-34 (HA0659) >gi 631472 pir  S4 5142 translation initiation factor eIF-4A2 homolog - human >gi 496902	2e-010	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4800	M76425	H.sapiens intron 2 Alu repetitive element.	0.014	<none></none>	<none></none>	<none></none>	
4801	X87212	H.sapiens mRNA for cathepsin C	0	1582221	prepro-cathepsin C [Homo sapiens]	1e-052	
4802	D80005	Human mRNA for KIAA0183 gene, partial cds	e-114	1136426	(D80005) KIAA0183 [Homo sapiens]	7e-025	
4803	AF026029	Homo sapiens poly(A) binding protein II (PABP2) gene, complete cds	2e-055	<none></none>	<none></none>	<none></none>	
4804	Z68322	Human DNA sequence from cosmid L79F5, Huntington's Disease Region, chromosome 4p16.3	2e-016	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.6	
4805	M63180	Human threonyl- tRNA synthetase mRNA, complete cds	0	135177	THREONYL- TRNA SYNTHETASE, CYTOPLASMIC (THREONINE TRNA LIGASE) (THRRS) 6.1.1.3) - human >gi 1464742 (M63180) threonyl-tRNA synthetase [Homo sapiens]	5e-070	
4806	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3.7	<none></none>	<none></none>	<none></none>	
4807	D16431	Human mRNA for hepatoma- derived growth factor, complete cds	3e-010	<none></none>	<none></none>	<none></none>	
4808	AF086168	Homo sapiens full length insert cDNA clone ZB82D09	e-148	1465826	(U64856) weak similarity to TPR domains [Caenorhabditis elegans]	2e-014	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4809	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.4	
4810	M34651	Pseudorabies virus with upstream and downsteam sequences.	0.4	417134	HEPATOCYTE NUCLEAR FACTOR 3-BETA norvegicus]	0.047	
4811	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	3e-010	1353390	(U34998) Rad9 [Coprinus cinereus]	3e-010	
4812	M94314	Homo sapiens ribosomal protein L30 mRNA, complete cds	1e-064	<none></none>	<none></none>	<none></none>	
4813	X95276	P.falciparum complete gene map of plastid- like DNA (IR-B)	0.001	<none></none>	<none></none>	<none></none>	
4814	X12716	Human Retrovirus mRNA for LTR (clone cH6)	5e-024	<none></none>	<none></none>	<none></none>	
4815	J03537	Human ribosomal protein S6 mRNA, complete cds.	e-138	133978	RIBOSOMAL PROTEIN S6 protein S6 - rat >gi 70933 pir  R3 MS6 ribosomal protein S6 - mouse >gi 319910 pir  R3 HU6 ribosomal protein S6 - human >gi 36148 (X67309) ribosomal protein S6 [Homo sapiens] >gi 54010 (Y00348) ribosomal protein S6 [Mus musculus] >g	3e-033	
4816	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4817	U61945	Caenorhabditis elegans cosmid C49C8.	1.8	<none></none>	<none></none>	<none></none>	
4818	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4819	M20020	Human ribosomal protein S6 mRNA, complete cds.	7e-072	225901	ribosomal protein S6 [Rattus norvegicus]	2e-015	
4820	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.058	<none></none>	<none></none>	<none></none>	
4821	AL023973	Human DNA sequence from clone 1033E15 on chromosome 22q13.1-13.2. Contains part of a novel gene, ESTs and a GSS, complete sequence [Homo sapiens]	3e-009	2352260	(AF000949) keratin [Canis familiaris]	0.037	
4822	M37430	Pea Chloroplast 4.5S, 5S, 16S and 23S mRNA.	4.7	4093193	(AF106583) unknown [Caenorhabditis elegans]	4.8	
4823	M63488	Human replication protein A 70kDa subunit mRNA complete cds.	0	1350579	REPLICATION PROTEIN A 70 KD DNA- BINDING SUBUNIT (RP-A) (RF-A) (REPLICATION FACTOR-A PROTEIN 1) (SINGLE- STRANDED DNA-BINDING PROTEIN) subunit [Homo sapiens]	8e-079	
4824	X83791	C.tentans BR1 gene	1.2	<none></none>	<none></none>	<none></none>	
4825	U67576	Methanococcus jannaschii section 118 of 150 of the complete genome	4	<none></none>	<none></none>	<none></none>	

		learest Neighbor		Nearest Neighbor			
	, , , , ,	astN vs. Genbank)	D 111 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	(BlastX vs. Non-Redundant Proteins)  ACCESSION   DESCRIPTION   P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE				
4826	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-009	<none></none>	<none></none>	<none></none>	
4827	X65319	Cloning vector pCAT-Enhancer	2e-077	987050	(X65335) lacZ gene product [unidentified cloning vector]	2e-011	
4828	X03558	Human mRNA for elongation factor 1 alpha subunit	0	1169475	ELONGATION FACTOR 1- ALPHA 1	e-109	
4829	X76538	H.sapiens Mpv17 mRNA	6.00E-98	730059	MPV17 PROTEIN >gi 631208 pir  S4 5343 glomerulosclerosis protein Mpv17 - human	3e-010	
4830	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4831	<none></none>	<none></none>	<none></none>	2078483	(U43200) antifreeze glycopeptide AFGP polyprotein precursor [Boreogadus saida]	0.014	
4832	X83617	H.sapiens mRNA for RanBP1	3.4	3924670	(AC004990)- supported by Genscan and several ESTs: C83049	3e-040	
4833	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	3024677	ISOLEUCYL- TRNA SYNTHETASE isoleucyl-tRNA synthetase (ileS) [Helicobacter pylori]	0.005	
4834	J02763	Human calcyclin gene, complete cds.	1e-043	<none></none>	<none></none>	<none></none>	
4835	L10910	Homo sapiens splicing factor (CC1.3) mRNA, complete cds.	0.00E+00	<none></none>	<none></none>	<none></none>	
4836	X53586	Human mRNA for integrin alpha 6	2e-099	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5	

	N	learest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4837	Z57594	H.sapiens CpG DNA, clone 186c5, reverse read cpg186c5.rt1b.	1.4	<none></none>	<none></none>	<none></none>	
4838	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4839	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
4840	Y00371	Human hsc70 gene for 71 kd heat shock cognate protein > :: gb AR013986 AR 013986 Sequence 15 from patent US 5773245	e-145	987050	(X65335) lacZ gene product [unidentified cloning vector]	7e-011	
4841	AF074991	Homo sapiens full length insert cDNA YH88A03	0.0005	<none></none>	<none></none>	<none></none>	
4842	AF055030	Homo sapiens clone 24538 mRNA sequence	2e-049	2842711	ZINC-FINGER PROTEIN UBI- D4 sapiens]	2e-016	
4843	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	1353531	(U38906) ORF14 [Bacteriophage r1t]	7.1	
4844	Z57588	H.sapiens CpG DNA, clone 186b7, reverse read cpg186b7.rt1b.	0.41	<none></none>	<none></none>	<none></none>	
4845	X65319	Cloning vector pCAT-Enhancer	9e-051	987050	(X65335) lacZ gene product [unidentified cloning vector]	0.37	
4846	X78411	B.pasteurii ureA, ureB and ureC genes.	3.1	<none></none>	<none></none>	<none></none>	
4847	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-009	2224697	(AB002376) KIAA0378 [Homo sapiens]	5e-008	

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)  ACCESSION   DESCRIPTION   P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION			
4848	U78729	Homo sapiens mad protein homolog Smad2 gene, exon 6	4.7	<none></none>	<none></none>	<none></none>	
4849	D55696	Human mRNA for cysteine protease, complete cds	0	2842759	LEGUMAIN PRECURSOR (ASPARAGINYL ENDOPEPTIDAS E) >gi 1743266 gnl PI D e286211 (Y09862) legumain [Homo sapiens]	3e-030	
4850	U95097	Xenopus laevis mitotic phosphoprotein 43 mRNA, partial cds	0.43	3005603	(AF053141) progesterone receptor [Equus caballus]	2.2	
4851	U46118	Rattus norvegicus cytochrome P450 3A9 mRNA, complete cds	0.38	<none></none>	<none></none>	<none></none>	
4852	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-006	2495726	HYPOTHETICAL PROTEIN KIAA0254 sapiens]	1e-005	
4853	L10911	Homo sapiens splicing factor (CC1.4) mRNA, complete cds.	e-117	<none></none>	<none></none>	<none></none>	
4854	D00132	Acremonium chrysogenum ARS DNA fragment	1.7	130998	SALIVARY PROLINE-RICH PROTEIN PRECURSOR (CLONE CP7) [CONTAINS: BASIC PEPTIDE P-F] glycoprotein precursor PRB2 - human (fragment) precursor [Homo sapiens]	0.45	
4855	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.9	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION		
4856	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4857	AC002186	Homo sapiens (subclone 1_f12 from P1 H115) DNA sequence	1e-041	2072966	(U93570) p40 [Homo sapiens]	4e-013	
4858	AF053520	Homo sapiens allele 12 fragile site locus	0.61	<none></none>	<none></none>	<none></none>	
4859	X65319	Cloning vector pCAT-Enhancer	2e-077	987050	(X65335) lacZ gene product [unidentified cloning vector]	2e-011	
4860	AJ005866	Homo sapiens mRNA for putative Sqv-7-like protein, partial	e-179	4008517	(AJ005866) Sqv- 7-like protein [Homo sapiens]	3e-049	
4861	AF052165	Homo sapiens clone 24522 mRNA sequence	4e-072	2065177	(Y12790) Supt5h protein [Homo sapiens] sapiens]	1e-021	
4862	M90058	Human serglycin gene, exons 1,2, and 3.	0.005	<none></none>	<none></none>	<none></none>	
4863	U17662	Human neurofibromatosis 1 (NF1) gene, exons 4c and 5 and partial cds	1.3	<none></none>	<none></none>	<none></none>	
4864	U64453	Human ELK1 pseudogene (ELK2) and immunoglobulin heavy chain gamma pseudogene (IGHGP)	3e-018	<none></none>	<none></none>	<none></none>	
4865	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
4866	X16826	Drosophila melanogaster DNA for 60C beta tubulin gene making beta 3 tubulin isoform	2.2	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID 1967	1105004	Vananus lasvis	4e-009	<none></none>	<none></none>	<none></none>	
4867	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	46-009				
4868	X65319	Cloning vector pCAT-Enhancer	8e-081	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	
4869	AL031322	S.pombe chromosome II cosmid c17D1	0.38	<none></none>	<none></none>	<none></none>	
4870	M11560	Human aldolase A mRNA, complete cds.	0	553861	(J05517) aldolase A [Mus musculus]	2e-066	
4871	U28831	Human protein immuno-reactive with anti-PTH polyclonal antibodies mRNA, partial cds. > :: gb I40055 I40055 Sequence 1 from patent US 5618695	e-106	896065	(U28831) protein that is immuno- reactive with anti- PTH polyclonal antibodies [Homo sapiens]	1e-014	
4872	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
4873	<none></none>	<none></none>	<none></none>		mucin, tracheal (AMN-22) - human (fragment)	4e-009	
4874	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds		<none></none>	<none></none>	<none></none>	
4875	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)	DALATEE	(BlastX vs. Non-Redundant Proteins)  ACCESSION   DESCRIPTION   P VALUE			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE				
4876	D85752	Enterococcus faecalis plasmid pPD1 bacA, bacB, bacC, bacD, bacE, bacF, bacG, bacH and bacI genes, complete cds	0.042	1123087	(U42436) C49H3.3 gene product [Caenorhabditis elegans]	0.001	
4877	AC001443	Homo sapiens (subclone 2_f10 from BAC 2913	1e-033	2072961	(U93568) putative p150 [Homo sapiens]	3e-007	
4878	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-012	<none></none>	<none></none>	<none></none>	
4879	S81433	heme oxygenase- 2 {5' region, alternative splicing}	4.2	<none></none>	<none></none>	<none></none>	
4880	M34312	S.cerevisiae telomeric sequence DNA, clone YLP108CA-4-ii.	5e-010	188864	(M74027) mucin [Homo sapiens]	2e-007	
4881	AF075079	Homo sapiens full length insert cDNA YQ80A08	1.00E-12	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	4.6	
4882	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.015	3176689	(AC003671) Contains similarity to ubiquitin carboxyl-terminal hydrolase 14 gb Z35927 from S. cerevisiae. [Arabidopsis thaliana]	4.5	
4883	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.12	<none></none>	<none></none>	<none></none>	
4884	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
H.	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4885	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
4886	U74586	Rattus norvegicus double-stranded RNA specific adenosine deaminase (RED2) mRNA, complete cds	3.5	2828280	(AL021687) putative protein [Arabidopsis thaliana] >gi 2832633 gnl PI D e1249651 (AL021711) putative protein [Arabidopsis thaliana]	4e-008	
4887	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	5e-014	2497599	LAMININ BETA- 2 CHAIN PRECURSOR	5.4	
4888	D78572	House mouse; Musculus domesticus mRNA for membrane glycoprotein, complete cds > :: dbj E12950 E129 50 cDNA GA3- 43 encoding novel polypeptide which appear when differentiate from embryo-tumor cell P19 to nerve cell	7e-017	1545807	(D78572) membrane glycoprotein [Mus musculus]	1.2	
4889	L07273	Rattus norvegicus carboxypeptidase E (CPE) gene, exon 1.	3.2	<none></none>	<none></none>	<none></none>	
4890	Z46629	Homo sapiens SOX9 mRNA. > :: gb G28593 G285 93 human STS SHGC-35378.	e-132	<none></none>	<none></none>	<none></none>	

	N	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4891	M30802	Human aromatase cytochrome P-450 gene, exon 8.	3.3	<none></none>	<none></none>	<none></none>	
4892	M28699	Homo sapiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds.	5e-088	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.2	
4893	M89955	Human 5-HT1D- type serotonin receptor gene, complete cds.	0	2494923	5- HYDROXYTRYP TAMINE 1D RECEPTOR 1D [Cavia porcellus]	3e-008	
4894	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
4895	AF004230	Homo sapiens monocyte/macrop hage Ig-related receptor MIR-7 (MIR cl-7) mRNA, complete cds	2e-012	<none></none>	<none></none>	<none></none>	
4896	D50463	Mouse SDR1 mRNA, complete cds	0	1806276	(X99337) glycoprotein 55 [Rattus norvegicus]	e-103	
4897	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4898	AB014597	Homo sapiens mRNA for KIAA0697 protein, partial cds	2e-067	3327208	(AB014597) KIAA0697 protein [Homo sapiens]	9e-051	
4899	AF047598	Homo sapiens origin recognition complex subunit 4 (ORC4L) mRNA, complete cds	e-110	2736149	(AF022108) putative replication initiator origin recognition complex subunit Orc4Lp [Homo sapiens] subunit 4; Orc4p [Homo sapiens]	7e-005	

	<u>N</u>	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID						316315	
4900	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA,	7e-007	<none></none>	<none></none>	<none></none>	
		complete cds					
4901	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
4902	U22325	Mus musculus faciogenital dysplasis (Fgd1) mRNA, complete cds.	1.20E+00	<none></none>	<none></none>	<none></none>	
4903	U22325	Mus musculus faciogenital dysplasis (Fgd1) mRNA, complete cds.	1.20E+00	<none></none>	<none></none>	<none></none>	
4904	U22325	Mus musculus faciogenital dysplasis (Fgd1) mRNA, complete cds.	1.20E+00	<none></none>	<none></none>	<none></none>	
4905	U26162	Human myosin regulatory light chain mRNA, complete cds.	0	228542	myosin:SUBUNIT =regulatory light chain	3e-068	
4906	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4907	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	3822225	(AF079183) RING-H2 finger protein RHG1a [Arabidopsis thaliana]	4e-006	
4908	X65319	Cloning vector pCAT-Enhancer	1e-075	987050	(X65335) lacZ gene product [unidentified cloning vector]	8e-019	
4909	AJ010475	Arabidopsis thaliana mRNA for DEAD box RNA helicase, RH28	0.62	<none></none>	<none></none>	<none></none>	

	N	learest Neighbor		Nearest Neighbor			
	(Bl	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4910	U48364	Mus musculus muscle-specific transcriptional activator alpha- NAC gp220 (Naca) mRNA, complete cds	0.2	<none></none>	<none></none>	<none></none>	
4911	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4912	J03750	Mouse single stranded DNA binding protein p9 mRNA, complete cds.	e-135	1709514	ACTIVATED RNA POLYMERASE II TRANSCRIPTIO NAL COACTIVATOR P15 (PC4) (P14) cofactor p15 - human >gi 531395 (U12979) PC4 [Homo sapiens] >gi 619161 (X79805) PC4, p15 [Homo sapiens]	1e-020	
4913	U70263	Border disease virus strain BD31, complete genome	3.2	<none></none>	<none></none>	<none></none>	
4914	AB012086	Canine herpesvirus gene for immediate- early protein, complete cds	0.37	<none></none>	<none></none>	<none></none>	
4915	X05908	Human mRNA for lipocortin	e-162	113944	ANNEXIN I (LIPOCORTIN I) (CALPACTIN II) (CHROMOBINDI N 9) (P35) (PHOSPHOLIPAS E A2 INHIBITORY PROTEIN) >gi 71756 pir  LU HU annexin I - human >gi 34388		
4916	<none></none>	<none></none>	<none></none>	*	<none></none>	<none></none>	
4917		Human clone 23652 mRNA sequence	0.13	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	(Bl	astN vs. Genbank)		` .	s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4918	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	8e-007	<none></none>	<none></none>	<none></none>	
4919	X57830	H.sapiens serotonin 5-HT2 receptor mRNA > :: gb G28536 G285 36 human STS SHGC-31576.	4e-011	<none></none>	<none></none>	<none></none>	
4920	U67559	Methanococcus jannaschii section 101 of 150 of the complete genome	3.5	<none></none>	<none></none>	<none></none>	
4921	M20020	Human ribosomal protein S6 mRNA, complete cds.	0	133978	A0S RIBOSOMAL PROTEIN S6 protein S6 - rat >gi 70933 pir  R3 MS6 ribosomal protein S6 - mouse >gi 319910 pir  R3 HU6 ribosomal protein S6 - human >gi 36148 (X67309) ribosomal protein S6 [Homo sapiens] >gi 54010 (Y00348) ribosomal protein S6 [Mus musculus] >g	2e-072	
4922	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	<none></none>	<none></none>	<none></none>	
4923	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	6e-006	<none></none>	<none></none>	<none></none>	
4924	X76683	Plasmid vector pHM2 betalactamase gene	e-160	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	

	Nearest Neighbor			Nearest Neighbor			
	(BlastN vs. Genbank)			(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4925	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4926	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	0.002	<none></none>	<none></none>	<none></none>	
4927	D50369	Homo sapiens mRNA for low molecular mass ubiquinone- binding protein, complete cds	e-152	3024781	UBIQUINOL- CYTOCHROME C REDUCTASE COMPLEX UBIQUINONE- BINDING PROTEIN QP-C PROTEIN) (COMPLEX III SUBUNIT VII) >gi 2605590 (D50369) low molecular mass ubiquinone- binding protein [Homo sapiens]	6e-023	
4928	M63391	Human desmin gene, complete cds.	4e-013	<none></none>	<none></none>	<none></none>	
4929	D38417	Mouse mRNA for arylhydrocarbon receptor, complete cds	e-110	<none></none>	<none></none>	<none></none>	
4930	U38253	Rattus norvegicus initiation factor eIF-2B gamma subunit (eIF-2B gamma) mRNA, complete cds	e-175	2494312	TRANSLATION INITIATION FACTOR EIF-2B GAMMA SUBUNIT (EIF- 2B GDP-GTP EXCHANGE FACTOR) subunit [Rattus norvegicus]	4e-040	
4931	D38417	Mouse mRNA for arylhydrocarbon receptor, complete cds	e-110	<none></none>	<none></none>	<none></none>	
4932	U50767	Mus musculus alpha 1 type I collagen gene, partial cds and 3' flanking region.	1.2	<none></none>	<none></none>	<none></none>	
4933	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	-MOINE>	

4.	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4934	U86137	Mus musculus telomerase protein-1 mRNA, complete cds	1.70E-01	3327208	(AB014597) KIAA0697 protein [Homo sapiens]	9e-006	
4935	S57980	Crp1=cystatin-related protein-1 [rats, Genomic, 7673 nt]	0.041	<none></none>	<none></none>	<none></none>	
4936	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4937	AB012047	Arabidopsis thaliana gene for sulfate transporter, complete cds, clone:AST56	0.14	3915658	ATP- DEPENDENT RNA HELICASE A helicase II [Homo sapiens]	6.1	
4938	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>	
4939	AB018374	Mus musculus GARP34 mRNA, complete cds	3e-037	<none></none>	<none></none>	<none></none>	
4940	AF001498	Campylobacter jejuni polysaccharide biosynthesis protein homolog gene, partial cds, galactosyl transferase homolog, UDP-galactose phosphate transferase homolog, acetyl transferase homolog and aminotransferase homolog gen	3e-005	<none></none>	<none></none>	<none></none>	
4941	J04617	Human elongation factor EF-1-alpha gene, complete cds. > :: dbj E02629 E026 29 DNA of human polypeptide chain elongation factor-		<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
		1 alpha					
4942	Z54349	H.sapiens MN/CA9 GENE	2e-007	<none></none>	<none></none>	<none></none>	
4943	AF077374	Homo sapiens small proline-rich protein (SPRR3) gene, exons 1, 2, and 3 and complete cds	1.3	<none></none>	<none></none>	<none></none>	
4944	X59828	Human chromosome 22 flanking hypervariable simple repeat DNA (clone HZREP42)	0.0003	<none></none>	<none></none>	<none></none>	
4945	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1.00E-09	124180	TRANSCRIPTIO NAL REGULATOR IE63 human- herpesvirus 1 (strain 17) herpesvirus 1] >gi 221713 (D00374) immediate early transcriptional modulating protein IE63 (gene UL54) herpesvirus 1]	5.8	
4946	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1.00E-09	124180	TRANSCRIPTIO NAL REGULATOR IE63 human herpesvirus 1 (strain 17) herpesvirus 1] >gi 221713 (D00374) immediate early transcriptional	5.8	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
					modulating protein		
					IE63 (gene UL54)		
					herpesvirus 1]		
		-					
] .							
4947	X76683	Plasmid vector	8e-092	987050	(X65335) lacZ	3e-015	
		рНМ2			gene product		
		betalactamase			[unidentified		
İ		gene			cloning vector]		
4948	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4949	U95094	Xenopus laevis	2.00E-04	<none></none>	<none></none>	<none></none>	
		XL-INCENP					
		(XL-INCENP)					
		mRNA, complete					
1050	7/1/0000	cds	1.000.01	10/0/00	1 3/11	1.0	
4950	X16972	Drosophila	1.20E-01	1362688	morphogen Xhh precursor - African	1.9	
	:	melanogaster cecropin gene			clawed frog		
		cluster			>gi 790938		
		Cluster			(L39213)		
					morphogen		
					[Xenopus laevis]		
4951	U12022	Human	0	2072296	(U95098) mitotic	5.9	
		calmodulin			phosphoprotein 44		
		(CALM1) gene,			[Xenopus laevis]		
1		exons 2,3,4,5 and		-			
		6, and complete					
4052	X56536	cds Rabbit mRNA for	2.3	119110	EBNA-1	4e-018	
4952	A30330	pH regulatory	2.3	119110	NUCLEAR	46-019	
		protein (Na+/H+			PROTEIN		
		exchanger),			herpesvirus 4		
		partial			(strain B95-8)	ļ	
		-			>gi 1334880	ĺ	
					(V01555) BKRF1		
					encodes EBNA-1		
					protein, latent	1	
					cycle gene.	l	
]					[Human	ľ	
					herpesvirus 4]		

	Nearest Neighbor			Nearest Neighbor			
	` ` `	astN vs. Genbank)	<del> </del>	(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4953	AF037438	Homo sapiens short chain L-3- hydroxyacyl-CoA dehydrogenase (SCHAD) gene, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
4954	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	4e-012	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	3.4	
4955	AB000467	Homo sapiens mRNA, partial cds, clone:RES4- 25	2e-012	<none></none>	<none></none>	<none></none>	
4956	U31525	Human glycogenin mRNA, complete cds	0	1707996	GLYCOGENIN >gi 2135280 pir  J C4695 glycogenin glucosyltransferas e (EC 2.4.1.186) - human	5e-042	
4957	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4958	AF003836	Mesocricetus auratus isopentenyl diphosphate:dime thylallyl diphosphate isomerase mRNA, complete cds	1.30E+00	<none></none>	<none></none>	<none></none>	
4959	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4960	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4961	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4.90E-02	<none></none>			
4962	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4.90E-02	<none></none>	<none></none>	<none></none>	
4963	L32537	Homo sapiens (clone XP6G6B) mRNA, partial EST.	5.00E-03	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor		
	,	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
4964	L32537	Homo sapiens (clone XP6G6B) mRNA, partial EST.	5.00E-03	<none></none>	<none></none>	<none></none>
4965	X63787	T.thermophila gene for snRNA U3-2	0.41	<none></none>	<none></none>	<none></none>
4966	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>
4967	U27341	Bos taurus endothelin converting enzyme-2 Sequence 1 from patent US 5736376	7e-015	<none></none>	<none></none>	<none></none>
4968	U35114	Human apolipoprotein E (APOE) gene, hepatic control region HCR-2	9e-005	<none></none>	<none></none>	<none></none>
4969	M86374	Rat tropoelastin gene, intron 25 (partial).	0.13	<none></none>	<none></none>	<none></none>
4970	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>
4971	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>
4972	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	<none></none>	<none></none>	<none></none>
4973	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7.00E-07	<none></none>	<none></none>	<none></none>
4974	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7.00E-07	<none></none>	<none></none>	<none></none>

	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4975	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	7.00E-07	<none></none>	<none></none>	<none></none>	
4976	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.005	2995290	(AL022268) putative transmembrane transport protein [Streptomyces coelicolor]	1.50E-02	
4977	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.005	2995290	(AL022268) putative transmembrane transport protein [Streptomyces coelicolor]	1.50E-02	
4978	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.005	2995290	(AL022268) putative transmembrane transport protein [Streptomyces coelicolor]	1.50E-02	
4979	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2.00E-05	2983512	(AE000718) putative protein [Aquifex aeolicus]	2.2	
4980	X56536	Rabbit mRNA for pH regulatory protein (Na+/H+ exchanger), partial	2.3	119110	EBNA-1 NUCLEAR PROTEIN herpesvirus 4 (strain B95-8) >gi 1334880 (V01555) BKRF1 encodes EBNA-1 protein, latent cycle gene. [Human herpesvirus 4]	4e-018	
4981	Z11508	A.thaliana rpl15 gene for plastid ribosomal protein CL15	5.00E-03	3283910	(AF070638) unknown [Homo sapiens]	2.5	
4982	X95834	H.sapiens DNA sequence surrounding NotI site, clone NRLA143D	7e-070	1588365	signal peptidase:SUBUN IT=12kD [Homo sapiens]	1e-043,	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
4983	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	2e-007	4008081	(AF106835) putative DnaJ [Methylovorus sp. strain SS1]	3e-010	
4984	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4985	U43626	Human chromosome 15q11-q13 putative DNA replication origin in the g- aminobutyric acid receptor b3 and a5 gene cluster	2e-018	2197085	(AF003535) ORF2-like protein [Homo sapiens]	0.0002	
4986	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4987	D21272	Rice mRNA for ADP-glucose pyrophosphorylas e	1.1	1708084	EXOGLUCANAS E B PRECURSOR 1,4-beta- cellobiosidase (EC 3.2.1.91) precursor - Cellulomonas fimi >gi 790698 (L38827) beta-1,4- cellobiohydrolase [Cellulomonas fimi]	5.8	
4988	U59706	Gallus gallus alternatively spliced AMPA glutamate receptor, isoform GluR2 flop, (GluR2) mRNA, partial cds.	0.015	<none></none>	<none></none>	<none></none>	
4989	AF086033	Homo sapiens full length insert cDNA clone YW26E09	e-174	<none></none>	<none></none>	<none></none>	
4990	L31840	Rattus norvegicus nuclear pore complex protein NUP107 mRNA, complete cds.	e-179	1709212	NUCLEAR PORE COMPLEX PROTEIN NUP107	2e-083	

1	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4991	AF052144	Homo sapiens clone 24573 and 24786 mRNA sequences	e-170	1174415	SPIDROIN 2 (DRAGLINE SILK FIBROIN 2) >gi 345426 pir  A4 4112 spidroin 2, dragline silk fibroin - orb spider (Nephila clavipes) (fragment) clavipes]	4.8	
4992	M22406	Human intestinal mucin mRNA, partial cds, clone SMUC 42.	0.085	188864	(M74027) mucin [Homo sapiens]	1e-009	
4993	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
4994	U24697	Chironomus samoensis nanos homolog (Cs nos) gene, complete cds.	0.13	3880999	(AL021492) Y45F10D.11 [Caenorhabditis elegans]	7e-022	
4995	M64716	Human ribosomal protein S25 mRNA, complete cds.	4e-074	2943738	(AB011550) Drosophila Policomblike- related gene containing PHD fingers. [Mus musculus]	4e-011	
4996	X54326	H.sapiens mRNA for glutaminyl- tRNA synthetase	0	135104	MULTIFUNCTIO NAL AMINOACYL- TRNA SYNTHETASE (CONTAINS: GLUTAMYL- TRNA SYNTHETASE glutamyl-prolyl- tRNA synthetase - human >gi 31958	1e-088	
4997	Z12112	pWE15A cosmid vector DNA	2e-028	987050	(X65335) lacZ gene product [unidentified cloning vector]	2e-007	
4998	Z62939	H.sapiens CpG DNA, clone 75f1, forward read cpg75f1.ft1b.	3e-010	<none></none>	<none></none>	<none></none>	

		Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)			s. Non-Redundant Pr	,	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
4999	<none></none>	<none></none>	<none></none>	2134574	mucin - rhesus macaque (fragment) >gi 437055	5e-005	
5000	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	9e-009	<none></none>	<none></none>	<none></none>	
5001	Z93950	H.sapiens DNA; chromosome Y repeat regions	0.15	<none></none>	<none></none>	<none></none>	
5002	X64037	H.sapiens mRNA for RNA polymerase II associated protein RAP74	5e-056	<none></none>	<none></none>	<none></none>	
5003	M37583	Human histone (H2A.Z) mRNA, complete cds.	e-132	121994	HISTONE H2A.Z  >gi 89608 pir  S03 642 histone  H2A.Z - bovine  >gi 92380 pir  S03 644 histone  H2A.Z - rat  >gi 106267 pir  A3 5881 histone  H2A.Z - human sapiens] >gi 57808 (X52316) histone  H2A.Z (AA 1- 127) taurus]  >gi 184060 (M37583) histone (H2A.Z) [Homo sapien	2e-044	
5004	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	<none></none>	<none></none>	<none></none>	
5005	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	4e-011	<none></none>	<none></none>	<none></none>	
5006	M94764	Glycine max cv. Dare nodulin 26 gene fragment.	0.043	<none></none>	<none></none>	<none></none>	

	N	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5007	Z34287	B.subtilis (SO113) genomic DNA (5425bp)	1.2	<none></none>	<none></none>	<none></none>	
5008	X76683	Plasmid vector pHM2 betalactamase gene	6e-078	987050	(X65335) lacZ gene product [unidentified cloning vector]	2e-014	
5009	D17577	Mouse mRNA for kinesin-like protein (Kiflb), complete cds	e-109	2497524	KINESIN-LIKE PROTEIN KIF1B mouse >gi 407339 gnl PI D d1005029 (D17577) Kif1b [Mus musculus]	9e-041	
5010	X91192	H.sapiens PLC beta 3 gene (exon 1) and SOM172 gene (exon 1)	1e-096	3294231	(AJ223970) monomethyl transferase	3	
5011	D88271	Human (lambda) DNA for immunogloblin light chain	1e-021	<none></none>	<none></none>	<none></none>	
5012	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
5013	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
5014	AF052133	Homo sapiens clone 23970 mRNA sequence	0	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.9	
5015	M21731	Human lipocortin-V mRNA, complete cds.	e-169	999934	Annexin V (Lipocortin V, Endonexin Ii, Placental Anticoagulant Protein) Mutant With Glu 17 Replaced By Gly, Glu 78 Replaced By Gln (E17g,E78q) Complexed With Calcium	4e-005	

	Nearest Neighbor			Nearest Neighbor			
	,	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5016	M21731	Human lipocortin-V mRNA, complete cds.	e-169	999934	Annexin V (Lipocortin V, Endonexin Ii, Placental Anticoagulant Protein) Mutant With Glu 17 Replaced By Gly, Glu 78 Replaced By Gln (E17g,E78q) Complexed With Calcium	4e-005	
5017	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
5018	L44118	Homo sapiens proximal CMT1A-REP repeat	0.0005	<none></none>	<none></none>	<none></none>	
5019	Y16849	Bacillus sp. D3 xynA and abfA genes and ORF1	2e-015	<none></none>	<none></none>	<none></none>	
5020	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	465975	PUTATIVE ATP-DEPENDENT RNA HELICASE T26G10.1 IN CHROMOSOME III >gi 482102 pir  S4 0731 ATP-dependent RNA helicase homolog T26G10.1 - Caenorhabditis elegans >gi 3880293 gnl PI D e1349766 1397- 1495 which introduced stop codon at 3' splice; 5' splice looks v.	9e-005	
5021	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	

13.57	Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)			s. Non-Redundant Pr		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5022	U02455	Cloning vector rpDR2, complete sequence.	0.35	2132302	hypothetical protein YPR144c - yeast similarity near C-terminus to RNA Polymerase beta subunit (Swiss Prot. accession number P11213) and CCAAT-binding transcription factor (PIR accession number A36368) [Saccharomyces cerevisiae]	1e-031	
5023	X97999	H.sapiens mRNA for transcription factor IID, subunit TAFII55	0	3024690	TRANSCRIPTIO N INITIATION FACTOR TFIID 55 KD SUBUNIT (TAFII-55) (TAFII55) factor IID [Homo sapiens]	4e-083	
5024	X71642	M.musculus GEG-154 mRNA	3e-092	<none></none>	<none></none>	<none></none>	
5025	X71642	M.musculus GEG-154 mRNA	3e-092	<none></none>	<none></none>	<none></none>	
5026	AB018270	Homo sapiens mRNA for KIAA0727 protein, partial cds	4e-061	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	7.6	
5027	D90086	Human pyruvate dehydrogenase (EC 1.2.4.1) beta subunit gene, exons 1-10	4e-011	2143936	probable regulatory protein 322 - rat	7.7	
5028	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.002	<none></none>	<none></none>	<none></none>	
5029	X65319	Cloning vector pCAT-Enhancer	2e-081	987050	(X65335) lacZ gene product [unidentified cloning vector]	3e-015	
5030	<none></none>	<none></none>	<none></none>	188864	(M74027) mucin [Homo sapiens]	0.001	

	Nearest Neighbor			Nearest Neighbor			
	(B)	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5031	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	0.0002	3258141	(AP000007) 138aa long hypothetical protein [Pyrococcus horikoshii]	9.6	
5032	X98001	H.sapiens mRNA for geranylgeranyl transferase II	e-129	2506788	GERANYLGERA NYL TRANSFERASE TYPE II BETA SUBUNIT (RAB GERANYLGERA NYLTRANSFER ASE BETA SUBUNIT) (RAB GERANYL- GERANYL- GERANYLTRAN SFERASE BETA SUBUNIT) transferase II [Homo sapiens]	3e-026	
5033	U727 <b>8</b> 9	Human cosmid U197H5, complete sequence [Homo sapiens]	5e-023	<none></none>	<none></none>	<none></none>	
5034	U72789	Human cosmid U197H5, complete sequence [Homo sapiens]	5e-023	<none></none>	<none></none>	<none></none>	
5035	U19239	Choristoneura fumiferana entomopoxvirus spheroidin gene, complete cds, G4R gene, partial cds, and nucleoside triphosphate phosphohydrolase (NPH I) gene, partial cds	3.8	<none></none>	<none></none>	<none></none>	
5036	U95098	Xenopus laevis mitotic phosphoprotein 44 mRNA, partial cds	8e-009	2690166	(AE000788) B. burgdorferi predicted coding region BBK23	4	

		Nearest Neighbor			Nearest Neighbor			
		lastN vs. Genbank)			s. Non-Redundant P			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
5037	U66871	Human enhancer of rudimentary homolog mRNA, complete cds		2498336	ENHANCER OF RUDIMENTARY HOMOLOG homologous to DROER protein [Homo sapiens] >gi 1519519 sapiens]	6e-057		
5038	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
5039	X99728	H.sapiens NDUFV3 gene, exon 3	3e-092	2829450	NADH- UBIQUINONE OXIDOREDUCT ASE 9 KD SUBUNIT PRECURSOR (COMPLEX I- 9KD) (CI-9KD)	1e-015		
5040	X78730	M. musculus DNA for the flanking sequences of the hypothalamic GRH first exons	2	<none></none>	<none></none>	<none></none>		
5041	X84373	H.sapiens mRNA for nuclear factor RIP140 > :: gb G28540 G285 40 human STS SHGC-31616.	e-155	<none></none>	<none></none>	<none></none>		
5042	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
5043	X82272	Human endogenous retrovirus env mRNA	8e-081	1196429	(M14123) pol/env ORF (bases 3878- 8257) first start codon at 4172; Xxx; putative [Homo sapiens]	6e-058		
5044	AF029982	Mus musculus sarco(endo)plasm ic reticulum calcium ATPase (SERCA2) gene, promoter region, exons 1-3, and partial cds	0.003	3873550	(AL033534) serine-rich protein	0.018		
5045	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		
5046	Y12781	Homo sapiens mRNA for transducin (beta) like 1 protein	1e-084	3021409	(Y12781) transducin (beta) like 1 protein [Homo sapiens]	2e-064		

	N	Nearest Neighbor		Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5047	S63912	D10S102=FBRN P [human, fetal brain, mRNA, 3043 nt]	4e-084	<none></none>	<none></none>	<none></none>	
5048	X91192	H.sapiens PLC beta 3 gene (exon 1) and SOM172 gene (exon 1)	1e-096	3294231	(AJ223970) monomethyl transferase	3	
5049	X03558	Human mRNA for elongation factor 1 alpha subunit	0	1169475	ELONGATION FACTOR 1- ALPHA 1	e-108	
5050	L31783	Mus musculus uridine kinase mRNA, partial cds	3e-029	1718058	URIDINE KINASE (URIDINE MONOPHOSPHO KINASE) >gi 471981 (L31783) uridine kinase	4e-011	
5051	X75652	A.longa plastid genes for tRNAs, ribosomal protein, rRNA and elongation factor	1.3	<none></none>	<none></none>	<none></none>	
5052	Z93123	M.acuminata mRNA; clone pBAN UD75	1.1	<none></none>	<none></none>	<none></none>	
5053	D16901	Human HepG2 3' region cDNA, clone hmd2h05	1.5	<none></none>	<none></none>	<none></none>	
5054	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.7	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5055	AF043252	Homo sapiens mitochondrial outer membrane protein (Tom40) gene, nuclear gene encoding mitochondrial protein, exons 7, 8 and 9	e-106	3941342	(AF043250) mitochondrial outer membrane protein [Homo sapiens] >gi 3941347 (AF043253) mitochondrial outer membrane protein [Homo sapiens] >gi 4105703 (AF050154) D19S1177E [Homo sapiens]	6e-007	
5056	X66494	R.norvegicus CHOT1 mRNA	1e-012	1545807	(D78572) membrane glycoprotein [Mus musculus]	3e-007	
5057	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
5058	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-007	3513368	(AB017202) entactin-2 [Mus musculus]	3e-005	
5059	U77107	Fundulus lineolatus cytochrome b (cytb) gene, mitochondrial gene encoding mitochondrial protein, partial cds	0.37	3947877	(AL034382) putative mitosis and maintenance of ploidy protein [Schizosaccharom yces pombe]	7e-026	
5060	X52317	Human mRNA for histone H2A.Z	5e-014	<none></none>	<none></none>	<none></none>	
5061	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-008	<none></none>	<none></none>	<none></none>	
5062	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1.2	<none></none>	<none></none>	<none></none>	

	Nearest Neighbor			Nearest Neighbor			
	2	lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5063	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
5064	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	1e-011	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	1.5	
5065	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0002	<none></none>	<none></none>	<none></none>	
5066	X15943	Huamn calcitonin/alpha- CGRP gene	1e-012	1575563	(U66464) hematopoietic progenitor kinase [Homo sapiens]	5.6	
5067	AF001175	Homo sapiens ribonuclease P protein subunit p14 (Rpp14) mRNA, complete cds	0	4100563	(AF001175) ribonuclease P protein subunit p14 [Homo sapiens]	2e-032	
5068	L29260	Arabidopsis thaliana 1-amino- 1- cyclopropanecarb oxylate synthase (ACS5) gene, complete cds.	0.41	<none></none>	<none></none>	<none></none>	
5069	X57268	Mouse DNA for t-haplotype- specific elements (located in H-2 complex, ETn related)	1.2	<none></none>	<none></none>	<none></none>	
5070	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	1e-010	2072296	(U95098) mitotic phosphoprotein 44 [Xenopus laevis]	5.5	
5071	Y11896	M.musculus mRNA for Brx gene, partial	3e-018	2196874	(Y11896) BRX protein [Mus musculus]	3e-011	

		Nearest Neighbor			Nearest Neighbor			
	`	astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)				
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE		
5072	Y00711	Human mRNA for lactate dehydrogenase B (LDH-B)	0	126041	L-LACTATE DEHYDROGENA SE H CHAIN dehydrogenase B (AA 1 - 334) [Homo sapiens] >gi 1200083	e-102		
5073	AF065482	Homo sapiens sorting nexin 2 (SNX2) mRNA, complete cds	0	3152938	(AF065482) sorting nexin 2 [Homo sapiens]	3e-072		
5074	M86374	Rat tropoelastin gene, intron 25 (partial).	0.13	<none></none>	<none></none>	<none></none>		
5075	D50418	Mouse mRNA for AREC3, partial cds	6e-047	2495271	SKELETAL MUSCLE- SPECIFIC ARE BINDING PROTEIN AREC3 (HOMEOBOX PROTEIN SIX4) M18) - mouse >gi 1255626 gnl PI D d1009550 (D50416) AREC3	2e-006		
5076	D17448	Microcystis aeruginosa plasmid pMA2 DNA, complete genome sequence	0.13	<none></none>	<none></none>	<none></none>		
5077	M29548	Human elongation factor 1-alpha (EF1A) mRNA, partial cds.	e-166	1169475	ELONGATION FACTOR 1- ALPHA 1	6e-010		
5078	AF081496	Homo sapiens kinetochore protein BUB3 (BUB3) mRNA, complete cds	6e-044	2921873	(AF047472) spleen mitotic checkpoint BUB3 [Homo sapiens] protein BUB3 [Homo sapiens]	3e-006		
5079	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>		
5080	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>		

	<u> </u>	Nearest Neighbor		Nearest Neighbor			
		lastN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5081	M14123	Human endogenous retrovirus HERV- K10.	2e-065	1196429	(M14123) pol/env ORF (bases 3878- 8257) first start codon at 4172; Xxx; putative [Homo sapiens]	6e-037	
5082	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
5083	D30655	Homo sapiens mRNA for eukaryotic initiation factor 4AII, complete cds	0	673433	(X56953) protein synthesis initiation factor 4A [Mus musculus]	2e-092	
5084	X16869	Human mRNA for elongation factor 1-alpha (clone CEF4)	5e-045	3122072	ELONGATION FACTOR 1- ALPHA 1 chicken >gi 488468 (L00677) elongation factor 1 alpha	1e-009	
5085	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
5086	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
5087	U78310	Homo sapiens pescadillo mRNA, complete cds	e-122	2194203	(U78310) pescadillo [Homo sapiens]	9e-009	
5088	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
5089	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-005	<none></none>	<none></none>	<none></none>	
5090	U09368	Human zinc finger protein ZNF140	0	1731416	ZINC FINGER PROTEIN 140 human >gi 487787 (U09368) zinc finger protein ZNF140	2e-062	
5091	M98509	Human NFB genomic fragment.	1e-010	<none></none>	<none></none>	<none></none>	
5092	AB002322	Human mRNA for KIAA0324 gene, partial cds	e-130	2996650	(AC004493) KIAA0324 [Homo sapiens]	9e-018	
5093	AJ007670	Homo sapiens mRNA for LGMD2B protein	2e-014	403460	(L24521) transformation- related protein [Homo sapiens]	3.8	

	Nearest Neighbor			Nearest Neighbor			
		astN vs. Genbank)		(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5094	X16869	Human mRNA for elongation factor 1-alpha (clone CEF4)	0	181967	(M29548) elongation factor 1-alpha [Homo sapiens]	2e-036	
5095	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	
5096	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0005	<none></none>	<none></none>	<none></none>	
5097	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
5098	· U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	2e-006	<none></none>	<none></none>	<none></none>	
5099	U45421	Borrelia burgdorferi 2.9-1 locus, ORF 5-8, ORF-A-D, REP+, REP-, and lipoprotein (LP) genes, complete cds	0.014	3510605	(AF044267) gyrase subunit B [Chlamydia trachomatis].	3.4	
5100	L54057	Homo sapiens CLP mRNA, partial cds.	0	<none></none>	<none></none>	<none></none>	
5101	D14660	Human mRNA for KIAA0104 gene, complete cds	0	1350786	PUTATIVE 60S RIBOSOMAL PROTEIN sapiens] >gi 3947438 (AC005034) ribosomal protein- like	e-111	

	N	learest Neighbor		Nearest Neighbor			
	(BlastN vs. Genbank)			(BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5102	X78627	H.sapiens mRNA for translin.	0	1082873	translin - human >gi 607130 (X78627) translin [Homo sapiens] >gi 1586346 prf  2 203413A recombination hotspot-binding protein [Homo sapiens]	5e-068	
5103	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	0.0001	<none></none>	<none></none>	<none></none>	
5104	M12585	Mouse alpha-1 antitrypsin gene, segment 1.	2e-006	3873550	(AL033534) serine-rich protein	1.7	
5105	X52967	Human mRNA for ribosomal protein L7	0	423072	ribosomal protein L7 - human	7e-061	
5106	U95094	Xenopus laevis XL-INCENP (XL-INCENP) mRNA, complete cds	7e-007	<none></none>	<none></none>	<none></none>	

	Nearest Nei	ghbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
5107	X78722	M.musculus GLUT2 gene for glucose transporter	0.34	1685115	(U68754) putative transcription factor [Dictyostelium discoideum]	3.8
5108	AF002677	Dictyostelium discoideum DEAD-box RNA helicase	0.28	3293508	(AF069188) NADH dehydrogenase 1 [Ephedrus laevicollis]	0.81
5109	AB018263	Homo sapiens mRNA for KIAA0720 protein, partial cds	0.87	107240	oncogene 1 (tre-2 locus) (clone 210) - human	0.19
5110	AF017115	Homo sapiens cytochrome c oxidase subunit IV precursor (COX4) gene, nuclear gene encoding mitochondrial protein, complete cds	0.77	<none></none>	<none></none>	<none></none>
5111	AE001383	Plasmodium falciparum chromosome 2, section 20 of 73 of the complete sequence	0.15	2315754	(AF016681) No definition line found [Caenorhabditis elegans]	9.6
5112	D49577	Pig mRNA for rearranged T-cell receptor delta- chain/Vdelta1.14- Ddeltas-Jdelta1, partial cds	0.91	<none></none>	<none></none>	<none></none>
5113	U63810	Homo sapiens WD40 protein Ciao 1 mRNA, complete cds	0.0	3219331	(AC004020) Unknown gene product [Homo sapiens]	3e-92
5114	AF085858	Homo sapiens full length insert cDNA clone YN49B07	e-172	3329465	(AF064553) NSD1 protein [Mus musculus]	<b>8</b> e-54
5115	X01682	Mouse gene for cytochrome P3-450	0.026	1381394	(U40989) tat interactive protein [Homo sapiens]	4.0

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID	ACCESSION	Descide from	1 VILLUE	ricobosion	Descrient from	1 111202	
5116	AE001432	Plasmodium	1.5	3873713	(Z74026) cDNA	9e-11	
3110	112001.02	falciparum			EST yk452h4.3		
		chromosome 2,			comes from this		
		section 69 of 73			gene; cDNA EST		
		of the complete			yk452h4.5 comes		
		sequence			from this gene		
5117	U31973	Human	2.3	136976	PROTEIN UL87	8.1	
		phosphodiesteras			>gi 76594 pir  S09		
		e A' subunit			851 hypothetical		
		(PDE6C) mRNA,			protein UL87 -		
		complete cds. > ::			human		
		gb G28549 G285	-		cytomegalovirus		
		49 human STS			cytomegalovirus]		
5110	3/00010	SHGC-31657.	2.6	<none></none>	AIONES	AIONES	
5118	X02212	Chicken alpha-	2.6	<none></none>	<none></none>	<none></none>	
5119	AE000838	cardiac actin gene Methanobacteriu	0.89	765086	(D30786) feline	1.4	
3119	AEUUU030	m	0.69	703080	CD9 [Felis catus]	1.4	
		thermoautotrophi			CD9 [Fells catus]		
		cum from bases					
		494834 to 505698					
		(section 44 of					
		148) of the					
		complete genome					
5120	U89744	Rattus norvegicus	0.68	728850	GLUCOAMYLA	9e-06	
		putative cell			SE S1/S2		
		surface antigen			PRECURSOR		
		mRNA, complete			(GLUCAN 1,4- ALPHA-		
		cds			GLUCOSIDASE)		
					(1,4-ALPHA-D-		
					GLUCAN		
1					GLUCOHYDROL		
					ASE)		
					>gi 626156 pir  S4		
					8478 glucan 1,4-		
					alpha-glucosidase		
					(EC 3.2.1.3) -		
					yeast stal, len:		
					1367, CAI: 0.3,	]	
					AMYH_YEAST P08640		
-					GLUCOAMYLA		
					SE S1 (EC	ļ	
					3.2.1.3)	ļ	
					[Saccharomyc		
	· · · · · · · · · · · · · · · · · · ·	L		L			

	Nearest Nei	ghbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
ID						
5121	J04974	Human alpha-2 type XI collagen mRNA (COL11A2).	1.2	114887	BREAKPOINT CLUSTER REGION PROTEIN protein, splice form 1 - human >gi 29421 (X02596) bcr gene product [Homo sapiens]	9.4
5122	AL021806	Homo sapiens DNA sequence from PAC 779B17 on chromosome 22q13.1. Contains exon trap, complete sequence	0.046	2827756	EPHRIN TYPE-A RECEPTOR 1 PRECURSOR	1.9
5123	X68826	P.sativum mRNA for fructose 1,6 biphosphatase	0.95	1314248	(U24681) NADH:cytochrom e c reductase [synthetic construct]	2e-05
5124	M14431	Bacteriophage phi-29 gene-16 gene, complete cds.	0.035	<none></none>	<none></none>	<none></none>
5125	U17033	Human 180 kDa transmembrane PLA2 receptor mRNA, complete cds.	0.36	722372	(U23139) similar to beta transducin proteins containing TRP- ASP domains [Caenorhabditis elegans]	3e-0 <b>8</b>
5126	Z50202	P.vulgaris arc5-1 gene	0.007	1151256	(U43319) transmembrane receptor [Mus musculus]	0.13
5127	AF013711	Homo sapiens 22 kDa actin-binding protein	2e-10	<none></none>	<none></none>	<none></none>
5128	AF086324	Homo sapiens full length insert cDNA clone ZD53E07	5e-09	3318653	(U83192) post- synaptic density protein 95 [Homo sapiens]	0.001
5129	D90117	T. thermophila mRNA for citrate synthase (EC 4.1.3.7)	0.63	<none></none>	<none></none>	<none></none>

	Nearest Nei	ghbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
5130	D45105	Metschnikowia reukaufii 26S rRNA, partial sequence	0.78	<none></none>	<none></none>	<none></none>	
5131	D85088	Ectoplana limuli DNA for 18s ribosomal RNA	0.41	267408	PROBABLE DNA PACKAGING PROTEIN packaging protein [Human herpesvirus 4]	7.2	
5132	X89886	P.patens mRNA for 5- aminolevulinate dehydratase	0.41	3875246	(Z81490) similar to WD domain, Gbeta repeats (2 domains); cDNA EST EMBL:T00482 comes from this gene; cDNA EST EMBL:T00923 comes from this gene; cDNA EST yk449d4.3 comes from this gene; cDNA EST yk449d4.5 comes from this gene; cDNA EST yk449d4.5 comes from this gen	2e-22	
5133	AB014564	Homo sapiens mRNA for KIAA0664 protein, partial cds	0.0	2981221	(AF053091) eyelid [Drosophila melanogaster]	0.076	
5134	AE001403	Plasmodium falciparum chromosome 2, section 40 of 73 of the complete sequence	0.003	2495297	HYPOTHETICAL 26.3 KD HOMEOBOX PROTEIN C02F12.5 IN CHROMOSOME X > gi 1109893 (U41545) strong similarity to homeobox proteins; similar to inhibitor domain of tissue factor pathway inhibitor	3.7	

	Nearest Nei	ghbor (BlastN vs. C	enbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
5135	U92574	Fugu rubripes homeobox protein HOXB-1 (FrHOXB-1) gene, complete cds	0.54	<none></none>	<none></none>	<none></none>	
5136	U31118	Xenopus laevis cytoplasmic myosin II regulatory light chain mRNA, complete cds	0.26	3879530	(Z49130) cDNA EST yk486b9.3 comes from this gene; cDNA EST yk486b9.5 comes from this gene	8e-07	
5137	L49035	Gorilla gorilla ABC-transporter (TAP2) mRNA, complete cds	0.21	4007066	(AJ131571) X protein [Hepatitis B virus]	1.3	
5138	AF068628	Mus musculus DNA cytosine-5 methyltransferase 3B3 (Dnmt3b) mRNA, alternatively spliced, complete cds	4e-04	<none></none>	<none></none>	<none></none>	
5139	M64982	Human fibrinogen alpha chain gene, complete mRNAs.	0.062	<none></none>	<none></none>	<none></none>	
5140	M19262	Rat clathrin light chain (LCB3) mRNA, complete cds.	0.25	2088802	(AF003151) D1007.4 gene product [Caenorhabditis elegans]	0.012	
5141	X94947	L.esculentum mRNA for homeobox protein	3.7	2315770	(AF016683) K09F6.1 gene product [Caenorhabditis elegans]	0.096	
5142	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	<none></none>	
5143	M33782	Human TFEB protein mRNA, partial cds.	0.36	<none></none>	<none></none>	<none></none>	
5144	AB011098	Homo sapiens mRNA for KIAA0526 protein, complete cds	2e-07	2501115	TBX2 PROTEIN (T-BOX PROTEIN 2)	0.90	

	Nearest Nei	ghbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
5145	AF039029	Homo sapiens snurportin1 mRNA, complete cds	0.0	3834390	(AF039029) snurportin1 [Homo sapiens]	e-108
5146	U22970	Human interferon- inducible peptide (6-16) gene, complete cds	0.21	<none></none>	<none></none>	<none></none>
5147	D63880	Human mRNA for KIAA0159 gene, complete cds	2e-64	<none></none>	<none></none>	<none></none>
5148	AB011174	Homo sapiens mRNA for KIAA0602 protein, partial cds	e-164	3043728	(AB011174) KIAA0602 protein [Homo sapiens]	2e-53
5149	AF053551	Homo sapiens metaxin 2 (MTX2) mRNA, nuclear gene encoding mitochondrial protein, complete cds	0.0	3283049	(AF053551) metaxin 2 [Homo sapiens]	1e-76
5150	¥13382	Arabidopsis thaliana ferrochelatase-I gene and promoter sequence	0.012	<none></none>	<none></none>	<none></none>
5151	AF044854	Colias eurytheme large subunit ribosomal RNA gene, partial sequence; tRNA-Val gene, complete sequence; and small subunit ribosomal RNA gene, partial sequence, mitochondrial genes for mitochondrial RNAs	1.3	<none></none>	<none></none>	<none></none>

	Nearest Nei	ghbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)		
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
5152	AF005059	Toxoplasma gondii p97 mRNA, complete cds	0.90	2570049	(Y08701) Pinin [Mus musculus]	1.3
5153	D84307	Human mRNA for phosphoethanola mine cytidylyltransfera se, complete cds	0.013	<none></none>	<none></none>	<none></none>
5154	D38050	Aspen prxA3a gene for peroxidase, complete cds	0.018	1723942	HYPOTHETICAL 20.8 KD PROTEIN IN COX4-GTS1 INTERGENIC REGION >gi 2131614 pir  S 61134 hypothetical protein YGL183c - yeast (Saccharomyces cerevisiae) >gi 1143564 gnl PI D e199057 (X91489) putative HMG box [Saccharomyces cerevisiae]	0.39
5155	AL010208	Plasmodium falciparum DNA *** SEQUENCING IN PROGRESS *** from contig 3-103, complete sequence	0.13	1850115	(Z86089) fadD2 [Mycobacterium tuberculosis]	1.5
5156	U07807	Human metallothionein IV (MTIV) gene, complete cds.	0.004	<none></none>	<none></none>	<none></none>
5157	AF048991	Homo sapiens MutS homolog 5 (MSH5) gene, exons 13 through 25 and complete cds	0.001	3986756	(AF109905) NG23 [Mus musculus]	0.007

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5158	U39079	Schizosaccharom yces pombe ARS binding protein 1	0.50	<none></none>	<none></none>	<none></none>	
5159	X01706	Mouse intracisternal A- particle (IAP) gene 62 long terminal repeat (LTR)	0.41	2224713	(AB002384) KIAA0386 [Homo sapiens]	8e-04	
5160	AF030558	Rattus norvegicus phosphatidylinosi tol 5-phosphate 4- kinase gamma mRNA, complete cds	8e-13	<none></none>	<none></none>	<none></none>	
5161	L06453	Strongylocentrotu s purpuratus (clone C) high mobility group 1 protein (HMG1 homologue) gene, complete cds.	0.33	3914031	BETA- GALACTOSIDE SPECIFIC LECTIN I A CHAIN (MLA) (ML-I A) (RRNA N- GLYCOSIDASE)	0.087	
5162	Z68320	Caenorhabditis elegans cosmid W07A12, complete sequence [Caenorhabditis elegans]	0.28	2500558	PUTATIVE RIBONUCLEASE III (RNASE III) >gi 3876420 gnl PI D e1346063 (Z81070) similar to ribonuclease [Caenorhabditis elegans]	2e-25	
5163	U40397	Mus musculus serum amyloid A- 4 protein (Saa4) gene, complete cds		<none></none>	<none></none>	<none></none>	
5164	X00367	Chlamydomonas chloroplast DNA region with ARS element 03 (ARS = autonomously replicating sequence)	0.046	<none></none>	<none></none>	<none></none>	

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IPTION   P VALUE
NE> <none></none>
84) 8e-04
B6 [Homo
NE> <none></none>
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CLEASE
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NE> <none></none>
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NE> <none></none>

	Nearest Ne	ighbor (BlastN vs. C	Genbank)	Nearest Neig	hbor (BlastX vs. Nor Proteins)	n-Redundant
CEO	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE
SEQ ID		DESCRIPTION				PVALUE
5165	U43838	Glycine max choline kinase GmCK1p mRNA, complete cds	1.2	132918	RIBOSOMAL PROTEIN L35, CHLOROPLAST PRECURSOR (CL35) >gi 81486 pir  A36 107 ribosomal protein L35 precursor, chloroplast - spinach oleracea]	2.4
5166	U67590	Methanococcus jannaschii section 132 of 150 of the complete genome	0.097	<none></none>	<none></none>	<none></none>
5167	AB006787	Mus musculus mRNA for apoptosis signal- regulating kinase 1, complete cds	0.39	1263187	(U24215) HOMODA hydrolase [Pseudomonas putida] putida]	0.83
5168	U43567	Trypanosoma cruzi kinetoplast maxicircle DNA, clone TRCKPMAX	0.054	<none></none>	<none></none>	<none></none>
5169	U04706	Bos taurus 50 kDa protein (adp50) mRNA, complete cds.	0.0	2498104	ADRENAL MEDULLA 50 KD PROTEIN	8e-83
5170	L14684	Rattus norvegicus nuclear-encoded mitochondrial elongation factor G mRNA, complete cds.	e-137		ELONGATION FACTOR G, MITOCHONDRI AL PRECURSOR (MEF-G) >gi 543383 pir  S4 0780 translation elongation factor G, mitochondrial - rat >gi 310102	3e-59
5171		Human glucose- 6-phosphatase mRNA, complete cds. >	2e-04		GLUCOSE-6- PHOSPHATASE (G6PASE) 3.1.3.9) - human >gi 452444 (U01120) glucose- 6-phosphatase [Homo sapiens]	4e-12

	Nearest Ne	ighbor (BlastN vs. (	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
5172	D87671	Rat mRNA for TIP120, complete cds	e-144	1799570	(D87671) TIP120 [Rattus norvegicus]	3e-69	
5173	U22296	Rattus norvegicus casein kinase 1 gamma 1 isoform mRNA, complete cds	e-120	3024053	CASEIN KINASE I, GAMMA 1 ISOFORM kinase 1 gamma 1 isoform [Rattus norvegicus]	8e-54	
5174	Y07648	A.thaliana nit2 gene, nit1 gene and nit3 gene	0.007	2429486	(AF025464) No definition line found [Caenorhabditis elegans]	9.5	
5175	AB013721	Oryctolagus cuniculus mRNA for mitsugumin 23, complete cds	3e-91	3628745	(AB013721) mitsugumin 23 [Oryctolagus cuniculus]	0.006	
5176	M74069	Saccharomyces cerevisiae endochitinase (CTS1-1) gene, complete cds.	2.5	<none></none>	<none></none>	<none></none>	
5177	Z61469	H.sapiens CpG DNA, clone 52h1, forward read cpg52h1.ft1a	1e-77	1184072	(U40766) COL-1 [Meloidogyne incognita]	0.002	
5178	AF015043	Homo sapiens EH-binding protein mRNA, partial cds	0.0		APOLIPOPROTE IN C-IV PRECURSOR cluster E-C1-C2 linked gene [Mus musculus]	3.0	
5179		H.sapiens (clone pS2) sequence	3e-04		(AL031798) putative diphthine synthase	3e-23	
5180	X94768	H.sapiens RP3 gene (XLRP gene 3)	1e-05	<none></none>	<none></none>	<none></none>	
5181		M.musculus mRNA for RIP1 protein	0.48		synapsin Ib - human	3e-04	
5182		Human Ig J chain gene, exons 3 and 4.	0.036	<none></none>	<none></none>	<none></none>	

***	Nearest Ne	ighbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5183	M30773	Human calcineurin B mRNA, complete cds	0.002	3878494	(Z79602) predicted using Genefinder; Similarity to Yeast hypothetical protein YAE2 gene; cDNA EST EMBL:M88949 comes from this gene	3e-06	
5184	U08831	Human immunodeficienc y virus type 1, sample 019 from Thailand (E2TH019W.01di 1sCD), envelope glycoprotein c2v3 region (env) gene, partial cds.		<none></none>	<none></none>	<none></none>	
5185	Z98303	Human DNA sequence from BAC 140H19 on chromosome Xq24-25. Contains STS	0.005	<none></none>	<none></none>	<none></none>	
5186	AE000952	Archaeoglobus fulgidus section 155 of 172 of the complete genome	2e-07	3257245	(AP000003) 571aa long hypothetical oxaloacetate decarboxylase alpha chain [Pyrococcus horikoshii]	5e-08	
5187	L48476	Homo sapiens (subclone 3_e10 from P1 H21) DNA sequence.	2e-04		(Z72510) similarity to yeast UTR3 protein (Swiss Prot accession number P21374); cDNA EST EMBL:D72822 comes from this gene; cDNA EST EMBL:D75763 comes from this gene; cDNA EST yk274e3.3 comes from this gene; cDNA EST	0.19	

	Nearest Nei	ghbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
					yk274e3		
5188	U95102	Xenopus laevis mitotic phosphoprotein 90 mRNA, complete cds	3e-09	<none></none>	<none></none>	<none></none>	
5189	AF055022	Homo sapiens clone 24684 mRNA sequence	e-102	2708743	(AC003952) putative Tal-1-like reverse transcriptase	4.0	
5190	AJ009761	Homo sapiens mRNA for putative dimethyladenosin e transferase, partial	e-121	4050050	(AF102147) putative dimethyladenosine transferase [Homo sapiens]	8e-48	
5191	Y08238	H.pylori clpB gene	0.27	1572756	(U70848) - C43G2.1 gene product [Caenorhabditis elegans]	1e-21	
5192		<none></none>	<none></none>	2828280	(AL021687) putative protein [Arabidopsis thaliana] >gi 2832633 gnl PI D e1249651 (AL021711) putative protein [Arabidopsis thaliana]	9e-36	
5193	J00747	Rat insulin-I (ins- 1) gene.	6e-05	4154522	(AE001441) putative [Helicobacter pylori]	3.2	
5194	U64454	Human 3' of immunoglobulin heavy chain locus	0.83	281204	gene LF3 protein - human herpesvirus 4 virus]	0.069	

***	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant			
000	4 00000001	DESCRIPTION.	D 174 1 115	AGOEGGIONI	Proteins)	DALATAR	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5195	AB002383	Human mRNA for KIAA0385 gene, complete cds	8e-13	2498318	DXS6673E PROTEIN retardation candidate gene [Homo sapiens]	2e-24	
5196	M81840	Human NRL gene product mRNA, complete cds.	0.029	3875740	(Z81497) similar to mannosyloligosaccharide alpha-1, 2-mannosidase; cDNA EST EMBL:D67155 comes from this gene; cDNA EST EMBL:D64219 comes from this gene; cDNA EST yk260e12.3 comes from this gene; cDNA EST yk260e12.5 comes from this gene; cDNA EST yk260e12.5 comes f	6e-18	
5197	U12523	Rattus norvegicus ultraviolet B radiation- activated UV98 mRNA, partial sequence.	1e-10	3219914	HYPOTHETICAL 16.8 KD PROTEIN C30D10.04 IN CHROMOSOME II >gi 2276353 gnl PI D e330328 pombe]	2e-11	
5198	AB017026	Mus musculus mRNA for oxysterol-binding protein, complete cds	0.0	3551523	(AB017026) oxysterol-binding protein	e-120	
5199	U83981	Homo sapiens apoptosis associated protein (GADD34) mRNA, complete cds	e-119	3258618	(U83981) apoptosis associated protein [Homo sapiens]	7e-26	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant			
				Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
5200	U37580	Streptomyces	0.048	2459916	(AF005859)	0.18	
		coelicolor			anon2D7		
	:	phosphotyrosine			[Drosophila		
		protein			melanogaster]		
		phosphatase					
		(ptpA) gene,					
		putative cystathionine					
		gamma-lyase					
		(cysA) gene, and					
		LysR-like protein					
		gene, complete					
		cds					
5201	D00723	Human mRNA	3e-19	<none></none>	<none></none>	<none></none>	
3201	200,23	for hydrogen	35 15	1,01,12	110112	TONE	
		carrier protein, a					
		component of an					
		enzyme complex,					
į į		glycine synthase					
		(EC 2.1.2.10)					
5202	X89366	A.thaliana DNA	0.025	1209669	(U38810) CAGR1	0.008	
		for 40 kDa			[Homo sapiens]		
		protein gene			>gi 3098420		
					(AF040945)		
					homeotic regulator homolog MAB21		
					[Mus musculus]		
5203	AF067158	HIV-1 isolate	2,4	<none></none>	<none></none>	<none></none>	
0203	111 00,100	301905 from	2	110112	4101,12	410112	
		India, complete					
		genome					
5204	U09954	Human ribosomal	5e-37	<none></none>	<none></none>	<none></none>	
		protein L9 gene,				1	
		5' region and				1	
		complete cds.					
5205	AF029984	Lycopersicon	7e-37	4090943	(AF029984)	2e-49	
		esculentum COP1			COP1 homolog	i	
		homolog (COP1)			[Lycopersicon	l	
		mRNA, complete cds			esculentum]	ł	
5206	U43076	Mus musculus	2e-17	2655422	(AF035530)	2e-22	
3200	043070	cdc37 homolog	20-1/	2033422	CDC37 [Gallus	2 <del>0</del> -22	
		mRNA, complete			gallus]		
		cds			guitusj		
				1	<u> </u>		

	Nearest Ne	ighbor (BlastN vs. (	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION		P VALUE	
ID						1 111202	
5207	U07745	Lycopersicon esculentum biotin-containing subunit of methylcrotonyl- CoA carboxylase mRNA, partial cds.	4e-32	533707	(U12536) 3- methylcrotonyl- CoA carboxylase precursor	4e-49	
5208	X74465	Human papillomavirus type 10 genomic DNA	1.3	3879121	(Z70310) predicted using Genefinder; Similarity to Mouse ankyrin (PIR Acc. No. S37771); cDNA EST EMBL:T01923 comes from this gene; cDNA EST EMBL:D32335 comes from this gene; cDNA EST EMBL:D32723 comes from this gene; cDNA EST EMBL:D32723 comes from this gene; cDNA EST EMBL:D32723 comes from this	2e-56	
5209		A.evecta gene encoding blue- light photoreceptor, intron	0.14	2257939	(AF005665) properdin [Homo sapiens]	7.6	
5210		Human tyrosine kinase arg gene mRNA.	1.1		(U42841) short region of weak similarity to chicken limb deformity protein (PIR:S24286) [Caenorhabditis elegans]	0.61	
5211	]	H.sapiens CpG DNA, clone 187a10, reverse read cpg187a10.rt1a.	e-102		(L10409) fork head related protein [Mus musculus]	1e-16	

	Nearest Nei	ghbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5212	X85753	Homo sapiens mRNA for CDK8 protein kinase > :: emb A61243 A61 243 Sequence 1 from Patent WO9709432	6e-59	1171821	NADH- UBIQUINONE OXIDOREDUCT ASE CHAIN 5 >gi 559499 gnl PI D e1192548 (X54253) ND5 protein	9.5	
5213	U27341	Bos taurus endothelin converting enzyme-2 Sequence 1 from patent US 5736376	7e-61	2136744	endothelin converting enzyme-2 - bovine	3e-29	
5214	U63648	Mus musculus p160 myb- binding protein (P160) mRNA, complete cds	4e-58	2645205	(U63648) p160 myb-binding protein [Mus musculus]	2e-34	
5215	AF035940	Homo sapiens MAGOH mRNA, complete cds	e-140	2306969	(AF007860) xl- Mago [Xenopus laevis]	3e-76	
5216	X80045	O.aries mRNA for acetyl-CoA carboxylase	2e-54	542750	acetyl-CoA carboxylase (EC 6.4.1.2) - human sapiens] >gi 740964 prf  20 06242A Ac-CoA carboxylase	8e-10	
5217	Z46372	R.norvegicus RNA for DNA topoisomerase II.	e-134	3876360	(Z68315) Similarity to Human MAP kinase phosphatase-1 (SW:PTN7_HUM AN) [Caenorhabditis elegans]	3e-12	
5218	AF035940	Homo sapiens MAGOH mRNA, complete cds	e-143	2330011	(AF007862) mm- Mago [Mus musculus] >gi 2909828 (AF035939) similar to mago nashi [Mus musculus] >gi 2909830	7e-81	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5219	Z72521	Human DNA sequence from cosmid N29F4 on chromosome 22q11.2-qter contains STS	6e-04	<none></none>	<none></none>	<none></none>	
5220	S74340	{clone E572, estrogen induced gene} [rats, Sprague-Dawley, hypothalamus, mRNA Partial, 130 nt]	4e-29	<none></none>	<none></none>	<none></none>	
5221	AL008711	Human DNA sequence from PAC 390N22 on chromosome Xp22.2	0.33	1184707	(U40868) folylpolyglutamate synthetase [Homo sapiens]	7.9	
5222	AE000012	Mycoplasma pneumoniae section 12 of 63 of the complete genome	0.15	<none></none>	<none></none>	<none></none>	
5223	D78333	Human mRNA for testis-specific TCP20, complete cds	e-113	2501141	T-COMPLEX PROTEIN 1, ZETA-LIKE SUBUNIT (TCP- 1-ZETA-LIKE) (CCT-ZETA- LIKE) TCP20 [Homo sapiens]	2e-42	
5224	AF042333	Oryza sativa 24- methylene lophenol C24(1)methyltran sferase mRNA, complete cds	0.003	3883124	(AF082300) arabinogalactan- protein [Arabidopsis thaliana]	0.006	
5225	U15426	Human anonymous mRNA sequence with CCA repeat region.	4e-06	1123105	(U42438) similar to S. cerevisiae longevity- assurance protein 1 (SP:P38703) [Caenorhabditis elegans]	0.34	

	Nearest Nei	ghbor (BlastN vs. C	Genbank)	Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5226	AF052497	Homo sapiens clone B18 unknown mRNA	0.003	1144514	(U34781) Antho- LWamidII preprohormone [Anthopleura elegantissima] >gi 1586846 prf  2 204411A prepro- hormone	4.3	
5227	D86590	Zinnia elegans mRNA for cinnamyl alcohol dehydrogenase, partial cds	0.13	<none></none>	<none></none>	<none></none>	
5228	AF081144	Rattus norvegicus CL1AA mRNA, complete cds	5e-14	1718004	TEGUMENT PROTEIN UL49 HOMOLOG herpesvirus 1] >gi 995634 (Z54206) UL49 [Bovine herpesvirus 1] >gi 2653299 gnl PI D e1187295 (AJ004801) virion protein (tegument) [Bovine herpesvirus type 1.1]	1.4	
5229	M63016	Human X chromosome enhancer-like sequence.	6e-04	<none></none>	<none></none>	<none></none>	
5230	L24755	Mus musculus bone morphogenetic protein (Bmp-1) mRNA, complete cds.	1.2	<none></none>	<none></none>	<none></none>	
5231	<none></none>	<none></none>	<none></none>		(AL021687) putative protein [Arabidopsis thaliana] >gi 2832633 gnl PI D e1249651 (AL021711) putative protein [Arabidopsis thaliana]	9e-36	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5232	U27341	Bos taurus endothelin converting enzyme-2 Sequence 1 from patent US 5736376	1e-22	2136744	endothelin converting enzyme-2 - bovine	2e-09	
5233	M81840	Human NRL gene product mRNA, complete cds.	0.030	3875740	(Z81497) similar to mannosyloligosaccharide alpha-1, 2-mannosidase; cDNA EST EMBL:D67155 comes from this gene; cDNA EST EMBL:D64219 comes from this gene; cDNA EST yk260e12.3 comes from this gene; cDNA EST yk260e12.5 comes from this gene; cDNA EST yk260e12.5 comes f	6e-18	
5234	AJ000097	Homo sapiens mRNA for EYA1B gene	2.7	3395586	(AL031179) similarity to phosphomannomu tases [Schizosaccharom yces pombe]	6e-38	
5235	U30788	Rattus norvegicus Tclone4 mRNA	1e-68	3523162	(AF076292) TGF- beta/activin signal transducer FAST- 1p	1.4	
5236	U88964	Human HEM45 mRNA, complete cds	0.0	2062680	(U88964) HEM45 [Homo sapiens]	7e-77	
5237	AF061016	Homo sapiens UDP-glucose dehydrogenase (UGDH) mRNA, complete cds	0.0	3127127	(AF061016) UDP- glucose dehydrogenase [Homo sapiens] dehydrogenase [Homo sapiens]	5e-90	
5238	D43921	Mouse AZ1 mRNA for pre- acrosome localization protein, complete cds	3e-15	2137118	acrosomal protein AZ1 - mouse localization protein [Mus musculus]	0.007	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant Proteins)			
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
5239	AF056022	Homo sapiens p60 katanin mRNA, complete cds	0.0	3283072	(AF056022) p60 katanin [Homo sapiens]	2e-60	
5240	U77949	Human Cdc6- related protein (HsCDC6) mRNA, complete cds	1e-83	<none></none>	<none></none>	<none></none>	
5241	AJ005016	Homo sapiens mRNA for putative ABC transporter, partial	0.0	3005931	(AJ005016) ABC transporter [Homo sapiens]	3e-70	
5242	X56756	Sheep mRNA for tumor necrosis factor alpha	4.5	<none></none>	<none></none>	<none></none>	
5243	AF020833	Homo sapiens eukaryotic translation initiation factor 3 subunit (p42) mRNA, complete cds	0.0	2460200	(AF020833) eukaryotic translation initiation factor 3 subunit [Homo sapiens]	e-158	
5244	X69878	H.sapiens Flt4 mRNA for transmembrane tyrosine kinase	4e-43	<none></none>	<none></none>	<none></none>	
5245	M27826	Human endogenous retroviral protease mRNA, complete cds.	1e-66	<none></none>	<none></none>	<none></none>	
5246	U20285	Human Gps1 (GPS1) mRNA, complete cds	2e-54	644879	(U20285) Gps1 [Homo sapiens]	8e-20	
5247	AF049528	Homo sapiens huntingtin-interacting protein HYPA/FBP11 (HYPA) mRNA, partial cds	5e-75	3341990	(AF049528) huntingtin- interacting protein HYPA/FBP11	2e-20	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant			
GEO	A COEGGIONI	DECOMPTON	DVALLE	A COEGGION	Proteins)	DALATAR	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
	1100000	 	0.14	0.67440	THE POST TOTAL AT	1 00	
5248	U87277	Human splicing	0.14	267449	HYPOTHETICAL	1e-08	
1		factor SRp30c			12.5 KD		
		gene, exon 1			PROTEIN ZK637.2 IN		
					CHROMOSOME		
					III		
					>gi 102507 pir  S1		
					5787 hypothetical		
					protein 1 (cosmid		
					ZK637) –		
					Caenorhabditis		
					elegans		
					Genefinder;		
					cDNA EST		
					yk217b5.3 comes		
					from this gene;		
					cDNA EST		
					yk217b5.5 comes		
					from this gene;		
					cDNA EST		
					yk340g12.3		
5249	D16919	Human HepG2 3'	e-164	3152559	(AC002986)	2e-52	
		region cDNA,			Similarity to A.		
		clone hmd3e06			thaliana gene		
					product		
						]	
					ınananaj		
5250	AJ006064	Rattus norvegicus	e-142	3757680	(AJ006064)	5e-73	
		mRNA for			coronin-like		
		coronin-like			protein [Rattus	ĺ	
		protein			norvegicus]		
5251	AR011000	Mue mueaulue	10.19	2780752	(AB006607)	0.001	
3231	ADVIIVU		16-10			0.001	
				11			
[					ine Rilase	İ	
		complete cds				l	
5250	AJ006064 AB011000	Mus musculus mRNA for choline/ethanola mine kinase,	e-142	3757680 2780752	protein [Rattus	5e-73	

	Nearest Neighbor (BlastN vs. Genbank)			Nearest Neighbor (BlastX vs. Non-Redundant			
					Proteins)		
SEQ	ACCESSION	DESCRIPTION	P VALUE	ACCESSION	DESCRIPTION	P VALUE	
ID							
5252	X80169	M.musculus mRNA for 200 kD protein	0.0	1717793	PROTEIN TSG24 (MEIOTIC CHECK POINT REGULATOR) >gi 1083553 pir  A 55117 tsg24 protein - mouse	e-150	

**Table 3** Polynucleotides encoding gene products of a protein family or having a known functional domain(s).

KIIOWI	runctional domain(s).					
SEQ II	D Validation Sequence	Biological Activity (Profile)		t Stop	Score	Direction
3920	393.E10.sp6:148957	7tm_1	531	710	9520	for
2667	172.F10.sp6:133946	7tm_2	45	724	8708	rev
2758	177.C6.sp6:134733	7tm 2	41	697	9828	rev
2933	184.C7.sp6:135556	7tm 2	3	834	8987	for
3129	121.E12.sp6:131940	7tm 2	245	1324	9550	rev
3365	172.A7.sp6:133883	7tm 2	94	761	8743	rev
3418	123.F9.sp6:132333	7tm 2	203	585	8785	rev
3419	123.F9.sp6:132333	7tm 2	203	585	8785	rev
3597	394.G2.sp6:149165	7tm 2	73	793	9209	for
3648	370.C5.sp6:141726	7tm 2	76	770	9269	for
3686	370.B1.sp6:141710	7tm 2	89	662	8791	for
3695	368.A12.sp6:141322	7tm_2	121	719	9015	rev
3696	368.A12.sp6:141322	7tm_2	121	719	9015	rev
4172	219.C10.sp6:139007	7tm 2	46	774	11394	rev
4216	368.D11.sp6:141357	7tm 2	66	775	9384	rev
4228	368.A11.sp6:141321	7tm 2	7	1079	9097	for
4441	99.F7.sp6:131296	7tm 2	534	1265	10956	rev
4442	99.F7.sp6:131296	7tm 2	534	1265	10956	rev
4482	100.D2.sp6:131459	7tm 2	122	1404	9296	rev
4495	395.B12.sp6:149307	7tm_2	79	1432	10427	rev
4525	90.B4.sp6:130874	7tm 2	4	691	9435	for
4616	100.D5.sp6:131462	7tm_2	655	1349	9255	for
4653	100.D7.sp6:131464	7tm 2	357	1346	11461	rev
4654	100.D7.sp6:131464	7tm_2	357	1346	11461	rev
4658	100.H6.sp6:131511	7tm 2	119	1035	10001	rev
4659	100.G6.sp6:131499	7tm 2	363	1188	9901	rev
4660	100.F6.sp6:131487	7tm_2	50	1127	8799	for
4661	100.F6.sp6:131487	7tm_2	50	1127	8799	for
4710	367.H9.sp6:141210	7tm_2	143	1266	11883	rev
4755	370.F4.sp6:141761	7tm_2	78	704	8942	for
4856	367.H11.sp6:141212	7tm_2	176	1227	9975	rev
4885	123.E10.sp6:132322	7tm_2	210	691	9071	rev
4900	123.E10.sp6:132322	7tm_2	210	691	9071	rev
4901	123.E10.sp6:132322	7tm_2	210	691	9071	rev
2656	176.H11.sp6:134606	ANK	207	290	4450	for
2555	180.C9.sp6:135947	asp	156	670	6710	for
3632	368.H11.sp6:141405	asp	136	1226	6880	rev
4205	368.B5.sp6:141327	asp	309	806	6073	for
4251	369.D6.sp6:141546	asp	434	1332	6263	rev
4253	396.F9.sp6:149544	asp	97	1106	5999	rev
4261	216.G10.sp6:139247	asp	74	703	6188	rev
4365	122.H12.sp6:132168	asp	152	1040	6183	rev
4498	80.H6.sp6:130297	asp	61	418	5944	rev
4664	172.E5.sp6:133929	asp	219	976	6434	for
4718	185.D9.sp6:135762	asp	31	872	5944	rev
4733	185.D9.sp6:135762	asp	31	872	5944	rev
4746	176.B10.sp6:134533	asp	253	1446	6079	rev

4822 17.F.F.s.p6:134766 asp 0 894 6336 rev 4854 184.F11.sp6:135596 asp 61 737 6416 rev 48636 367.HI1.sp6:141212 asp 81 1187 6182 rev 4929 180.E6.sp6:135968 asp 81 706 6150 for 2842 217.HI1.sp6:139452 ATPases 135 627 11664 for 2842 217.HI1.sp6:139452 ATPases 170 616 6150 for 2842 217.HI1.sp6:139452 ATPases 170 616 6150 for 2842 217.HI1.sp6:139452 ATPases 170 616 6150 for 2843 ATPases 170 616 6150 for 2844	SEQ ID NO:	Validation Sequence	Biological Activity (Profile)	Start	Stop	Score	Direction
A856   367.H11.sp6:141212   asp   81   1187   6182   rev     4929   180.E6.sp6:135968   asp   81   706   6150   for     4931   180.E6.sp6:135968   asp   81   706   6150   for     4931   180.E6.sp6:135976   ATPases   135   627   11664   for     2723   180.F2.sp6:135976   ATPases   2   701   5972   for     3019   216.B1.sp6:1319178   ATPases   170   616   6150   for     3046   121.B8.sp6:131900   ATPases   13   635   5867   rev     3190   80.D2.sp6:130245   ATPases   13   386   6068   for     3290   176.C6.sp6:134541   ATPases   329   730   6206   for     3290   176.C6.sp6:149183   ATPases   329   730   6206   for     3998   394.H8.sp6:149183   ATPases   313   816   6057   for     4119   218.F11.sp6:138852   ATPases   313   816   6057   for     4159   219.A7.sp6:138552   ATPases   318   8662   6145   for     4223   368.F9.sp6:141379   ATPases   38   6622   6145   for     4384   181.G11.sp6:135354   ATPases   362   769   5900   rev     4473   369.B4.sp6:141520   ATPases   36   605   6001   rev     4468   367.H8.sp6:14910   ATPases   36   605   6001   rev     4689   367.H8.sp6:14535578   ATPases   333   813   5773   for     4785   184.E5.sp6:135558   ATPases   333   813   5773   for     4847   184.B11.sp6:135554   ATPases   333   813   5773   for     4847   184.B11.sp6:135558   ATPases   333   813   5773   for     4849   369.D3.sp6:141918   ATPases   4   655   5933   for     4840   176.F10.sp6:134581   Bcl-2   69   356   16419   for     3404   176.F10.sp6:134581   Bcl-2   69   356   16419   for     3408   172.E1.sp6:133993   BZIP   116   304   5931   for     3408   172.E1.sp6:133995   BZIP   116   304   5931   for     3408   172.E1.sp6:133991   Cys-protease   38   992   10103   rev     4850   172.E9.sp6:134961   Cys-protease   38   992   10103   rev     4868   180.D1.sp6:133991   Cys-protease   38   992   10103   rev     4868   180.D1.sp6:133991   Cys-protease   38   992   10103   rev     4869   367.B7.sp6:141182   Dad_box_helic   Trev   S460   Trev     4871   377.As.sp6:14195   Dad_box_helic   Trev   S460   Trev	4822	177.F3.sp6:134766		0	894	6336	rev
4929   180.E6.sp6:135968   asp   81   706   6150   for	4854	184.F11.sp6:135596	asp	61	737	6416	rev
4931   180.E6.sp6:135968   asp   81   706   6150   for	4856	367.H11.sp6:141212	asp	81	1187	6182	rev
2723         180.F2.sp6:135976         ATPases         135         627         11664         for           2842         217.H11.sp6:139452         ATPases         2         701         5972         for           3019         216.B1.sp6:139178         ATPases         170         616         6150         for           3190         80.D2.sp6:130245         ATPases         13         386         6068         for           3290         176.C6.sp6:134541         ATPases         35         587         583         for           3670         369.C10.sp6:141538         ATPases         329         730         6206         for           3998         394.H8.sp6:149183         ATPases         21         571         5954         rev           4159         219.A7.sp6:138890         ATPases         313         816         6057         for           4159         219.A7.sp6:1388980         ATPases         18         648         5937         for           4159         219.A7.sp6:138813         ATPases         18         662         6145         for           4223         368.Ps.p6:141520         ATPases         42         412         14130         for	4929	180.E6.sp6:135968	asp	81	706	6150	for
2842 217.H11.sp6:139452 ATPases 2 701 5972 for 3019 216.B1.sp6:139178 ATPases 170 616 6150 for 3046 121.B8.sp6:131900 ATPases 13 635 5867 rev 3190 80.D2.sp6:130245 ATPases 13 386 6068 for 3290 176.C6.sp6:134541 ATPases 85 579 5883 for 3670 369.C10.sp6:141538 ATPases 21 571 5954 rev 4119 218.F11.sp6:138852 ATPases 313 816 6057 for 4159 219.A7.sp6:138980 ATPases 178 648 5937 for 423 368.F9.sp6:141379 ATPases 178 648 5937 for 4384 181.G11.sp6:135354 ATPases 362 769 5900 rev 4473 369.B4.sp6:141520 ATPases 4 412 14130 for 4540 218.C8.sp6:138813 ATPases 12 576 5782 rev 4689 367.H8.sp6:141209 ATPases 17 476 5905 rev 4785 184.E5.sp6:135578 ATPases 184 632 5943 for 4792 184.C6.sp6:135555 ATPases 333 813 5773 for 4847 184.B11.sp6:135548 ATPases 4 655 5933 for 4792 184.C6.sp6:135554 ATPases 4 655 5933 for 4792 184.C5.sp6:141918 ATPases 4 655 5933 for 4792 184.C5.sp6:14182 bromodomain 40 210 8810 for 4869 369.D3.sp6:141918 ATPases 4 655 5933 for 4792 184.C5.sp6:135939 BZIP 116 304 5931 for 4850 172.E1.sp6:133925 BZIP 146 298 4066 for 3951 393.G5.sp6:148976 BZIP 116 304 5931 for 4850 172.E1.sp6:133925 BZIP 116 304 5931 for 4850 172.E1.sp6:13393 BZIP 91 260 4366 for 5951 393.G5.sp6:14966 Cyclin 11 281 6930 for 4688 180.D1.sp6:135951 Cys-protease 72 348 184.79 for 4684 180.D1.sp6:135951 Cys-protease 72 348 184.79 for 4684 180.D1.sp6:135951 Cys-protease 72 348 184.79 for 4684 180.D1.sp6:135951 Cys-protease 72 348 184.79 for 4684 180.D1.sp6:135951 Cys-protease 73 370. 570 for 5830 185.A1.sp6:134799 EFhand 477 563 370. for 5830 185.A1.sp6:134799 EFhand 477 563 370. for 5830 185.A1.sp6:134799 EFhand 477 563 370. for 6830 185.A1.sp6:134799 EFhand 477 563 370. for 6830 185.A1.sp6:134799 EFhand 477 563 370. for 6830 185.A1.sp6:134799 EFhand 477 563 370. for 6830 185.A1.sp6:134803 EFhand 40 114 2640 rev 4154 218.E10.sp6:138803 EFhand 40 114 2640 rev 4154 218.E10.sp6:138803 EFhand 40 114 2640 rev 4154 218.E10.sp6:138803 EFhand 40 114 2640 rev 4154 218.E10.sp6:138803 EFhand 40 114 2640 rev 4154 218.E10.sp6:138803 EFhand 40 114 2640 rev	4931	180.E6.sp6:135968	asp	81	706	6150	for
3019   216.B1.sp6:139178   ATPases   170   616   6150   for	2723	180.F2.sp6:135976	ATPases	135	627	11664	for
3046         121.B8.sp6:131900         ATPases         13         635         5867         rev           3190         80.D2.sp6:130245         ATPases         13         386         6068         for           3290         176.C6.sp6:134541         ATPases         85         579         5883         for           369.C10.sp6:141538         ATPases         329         730         6206         for           3998         394.H8.sp6:149183         ATPases         21         571         5954         rev           4119         218.F1.sp6:138852         ATPases         313         816         6057         for           4159         219.A7.sp6:138980         ATPases         88         662         6145         for           4223         368.P9.sp6:141520         ATPases         178         648         5937         for           4384         181.G1.sp6:138813         ATPases         4         412         14130         for           4540         218.C8.sp6:138181         ATPases         12         576         5782         rev           4765         184.D5.sp6:135578         ATPases         17         476         5905         rev           4782 <td>2842</td> <td>217.H11.sp6:139452</td> <td>ATPases</td> <td>2</td> <td>701</td> <td>5972</td> <td>for</td>	2842	217.H11.sp6:139452	ATPases	2	701	5972	for
3190         80.D2.sp6:130245         ATPases         13         386         6068         for           3290         176.C6.sp6:134541         ATPases         85         579         5883         for           3670         369.C10.sp6:141538         ATPases         329         730         6206         for           3998         394.H8.sp6:149183         ATPases         21         571         5954         rev           4119         218.F11.sp6:138852         ATPases         313         816         6057         for           4159         219.A7.sp6:138852         ATPases         362         668         5937         for           4159         219.A7.sp6:138890         ATPases         362         769         5900         rev           4223         368.F9.sp6:141520         ATPases         362         769         5900         rev           4473         369.B4.sp6:135534         ATPases         4         412         14130         for           4560         404.G6.sp6:162933         ATPases         12         576         5782         rev           4785         184.E5.sp6:135558         ATPases         186         605         6001         rev	3019	216.B1.sp6:139178	ATPases	170	616	6150	for
3190         80.D2.sp6:130245         ATPases         13         386         6068         for           3290         176.C6.sp6:134541         ATPases         85         579         5883         for           3670         369.C10.sp6:141538         ATPases         329         730         6206         for           3998         394.H8.sp6:149183         ATPases         21         571         5954         rev           4119         218.F11.sp6:138852         ATPases         313         816         6057         for           4159         219.A7.sp6:138880         ATPases         88         662         6145         for           4223         368.F9.sp6:141379         ATPases         362         769         5900         rev           4473         369.B4.sp6:141520         ATPases         4         412         14130         for           4560         404.G6.sp6:162933         ATPases         86         605         6001         rev           4689         367.H8.sp6:141520         ATPases         18         632         5943         for           4785         184.E5.sp6:1355578         ATPases         18         605         6001         rev	3046	121.B8.sp6:131900	ATPases	13	635	5867	rev
3290         176.C6.sp6:134541         ATPases         85         579         5883         for           3670         369.C10.sp6:141538         ATPases         329         730         6206         for           3998         394.H8.sp6:149183         ATPases         21         571         5954         rev           4119         218.F11.sp6:138852         ATPases         313         816         6057         for           4159         219.A7.sp6:138980         ATPases         178         648         5937         for           4223         368.F9.sp6:141379         ATPases         178         648         5937         for           4384         181.G11.sp6:135354         ATPases         362         769         5900         rev           4473         369.B4.sp6:141520         ATPases         4         412         14130         for           4560         404.G6.sp6:162933         ATPases         86         605         6001         rev           4689         367.H8.sp6:141209         ATPases         17         476         5905         rev           4785         184.E5.sp6:135578         ATPases         184         632         5943         for	3190	80.D2.sp6:130245	ATPases	13	386	6068	for
3670         369.C10.sp6:141538         ATPases         329         730         6206         for           3998         394.H8.sp6:149183         ATPases         21         571         5954         rev           4119         218.F11.sp6:138852         ATPases         313         816         6057         for           4159         219.A7.sp6:138980         ATPases         88         662         6145         for           4223         368.F9.sp6:141379         ATPases         178         648         5937         for           4384         181.G11.sp6:135354         ATPases         362         769         5900         rev           4540         218.C8.sp6:135813         ATPases         4         412         14130         for           4560         404.G6.sp6:162933         ATPases         86         605         6001         rev           4785         184.E5.sp6:135578         ATPases         17         476         5905         rev           4785         184.E5.sp6:135558         ATPases         184         632         5943         for           4792         184.C6.sp6:135548         ATPases         14         498         6140         for	3290	<u>=</u>	ATPases	85	579	5883	for
3998         394.H8.sp6:149183         ATPases         21         571         5954         rev           4119         218.F11.sp6:138852         ATPases         313         816         6057         for           4159         219.A7.sp6:138980         ATPases         88         662         6145         for           4223         368.F9.sp6:141379         ATPases         178         648         5937         for           4384         181.G11.sp6:135354         ATPases         362         769         5900         rev           4473         369.B4.sp6:141520         ATPases         4         412         14130         for           4560         404.G6.sp6:162933         ATPases         6         605         6001         rev           4689         367.H8.sp6:141209         ATPases         17         476         5905         rev           4785         184.E5.sp6:135578         ATPases         184         632         5943         for           4787         184.B5.sp6:135554         ATPases         14         498         6140         for           4847         184.B11.sp6:13581         BCl-2         69         356         16419         for      <		<del>-</del>	ATPases	329	730	6206	for
4119         218.F11.sp6:138852         ATPases         313         816         6057         for           4159         219.A7.sp6:138980         ATPases         88         662         6145         for           4223         368.F9.sp6:141379         ATPases         178         648         5937         for           4384         181.G11.sp6:135354         ATPases         362         769         5900         rev           4473         369.B4.sp6:141520         ATPases         4         412         14130         for           4560         218.C8.sp6:138813         ATPases         12         576         5782         rev           4689         367.H8.sp6:141209         ATPases         16         605         6001         rev           4785         184.E5.sp6:135578         ATPases         184         632         5943         for           4792         184.C6.sp6:1355548         ATPases         14         498         6140         for           4847         184.B1.sp6:134581         Bcl-2         69         356         16419         for           4036         367.F5.sp6:14182         bromodomain         40         210         8810         for		-	ATPases	21		5954	rev
4159         219.A7.sp6:138980         ATPases         88         662         6145         for           4223         368.F9.sp6:141379         ATPases         178         648         5937         for           4384         181.G11.sp6:135354         ATPases         362         769         5900         rev           4473         369.B4.sp6:141520         ATPases         4         412         14130         for           4540         218.C8.sp6:138813         ATPases         12         576         5782         rev           4560         404.G6.sp6:162933         ATPases         86         605         6001         rev           4689         367.H8.sp6:141209         ATPases         184         632         5943         for           4785         184.E5.sp6:1355578         ATPases         184         632         5943         for           4782         184.B5.sp6:135554         ATPases         184         632         5943         for           4792         184.C6.sp6:134581         ATPases         14         498         6140         for           491         377.C1.sp6:141818         ATPases         4         655         5933         for		<u>.</u>	ATPases		816	6057	
4223       368.F9.sp6:141379       ATPases       178       648       5937       for         4384       181.G11.sp6:135354       ATPases       362       769       5900       rev         4473       369.B4.sp6:141520       ATPases       4       412       14130       for         4540       218.C8.sp6:138813       ATPases       12       576       5782       rev         4560       404.G6.sp6:162933       ATPases       86       605       6001       rev         4689       367.H8.sp6:141209       ATPases       17       476       5905       rev         4785       184.E5.sp6:135578       ATPases       184       632       5943       for         4792       184.C6.sp6:135555       ATPases       333       813       5773       for         4847       184.B11.sp6:135548       ATPases       14       498       6140       for         4036       367.F5.sp6:134918       BCl-2       69       356       16419       for         4489       369.D3.sp6:141543       bromodomain       63       230       10270       for         4489       369.D3.sp6:1349876       BZIP       146       298       4066		_	ATPases	88		6145	for
4384         181.G11.sp6:135354         ATPases         362         769         5900         rev           4473         369.B4.sp6:141520         ATPases         4         412         14130         for           4540         218.C8.sp6:138813         ATPases         12         576         5782         rev           4560         404.G6.sp6:162933         ATPases         86         605         6001         rev           4560         404.G6.sp6:141209         ATPases         17         476         5905         rev           4689         367.H8.sp6:1345578         ATPases         184         632         5943         for           4792         184.C6.sp6:135555         ATPases         184         632         5943         for           4847         184.B11.sp6:135548         ATPases         14         498         6140         for           5041         377.C1.sp6:141918         ATPases         4         655         5933         for           4489         369.D3.sp6:14182         bromodomain         40         210         8810         for           4489         369.D3.sp6:14182         bromodomain         63         230         10270         for <td></td> <td>-</td> <td>and the second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second 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second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second s</td> <td></td> <td></td> <td>5937</td> <td></td>		-	and the second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second second 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4540         218.C8.sp6:138813         ATPases         12         576         5782         rev           4560         404.G6.sp6:162933         ATPases         86         605         6001         rev           4689         367.H8.sp6:141209         ATPases         17         476         5905         rev           4785         184.E5.sp6:135578         ATPases         184         632         5943         for           4792         184.C6.sp6:135555         ATPases         333         813         5773         for           4847         184.B11.sp6:135548         ATPases         14         498         6140         for           5041         377.C1.sp6:141918         ATPases         4         655         5933         for           4036         367.F5.sp6:141543         Bcl-2         69         356         16419         for           4489         369.D3.sp6:141543         bromodomain         63         230         10270         for           4485         172.E1.sp6:133925         BZIP         146         298         4066         for           395.1         393.G5.sp6:148976         BZIP         116         304         5931         for <t< td=""><td></td><td><del>.</del></td><td></td><td></td><td></td><td></td><td></td></t<>		<del>.</del>					
4560         404.G6.sp6:162933         ATPases         86         605         6001         rev           4689         367.H8.sp6:141209         ATPases         17         476         5905         rev           4785         184.E5.sp6:135578         ATPases         184         632         5943         for           4792         184.C6.sp6:1355558         ATPases         333         813         5773         for           4847         184.B11.sp6:135548         ATPases         14         498         6140         for           5041         377.C1.sp6:141918         ATPases         4         655         5933         for           3404         176.F10.sp6:134581         Bcl-2         69         356         16419         for           4036         367.F5.sp6:141182         bromodomain         40         210         8810         for           4489         369.D3.sp6:148976         BZIP         146         298         4066         for           3951         393.G5.sp6:148976         BZIP         116         304         5931         for           4850         172.E9.sp6:133933         BZIP         91         260         4366         for		<del>-</del>					
4689         367.H8.sp6:141209         ATPases         17         476         5905         rev           4785         184.E5.sp6:135578         ATPases         184         632         5943         for           4792         184.C6.sp6:135555         ATPases         333         813         5773         for           4847         184.B11.sp6:135548         ATPases         14         498         6140         for           5041         377.C1.sp6:141918         ATPases         4         655         5933         for           3404         176.F10.sp6:134581         Bcl-2         69         356         16419         for           4036         367.F5.sp6:141182         bromodomain         40         210         8810         for           4489         369.D3.sp6:141543         bromodomain         63         230         10270         for           3408         172.E1.sp6:133925         BZIP         116         304         5931         for           4850         172.E9.sp6:133933         BZIP         116         304         5931         for           4850         370.B12.sp6:141721         cyclin         11         281         6930         for		-					
4785       184.E5.sp6:135578       ATPases       184       632       5943       for         4792       184.C6.sp6:135555       ATPases       333       813       5773       for         4847       184.B11.sp6:135548       ATPases       14       498       6140       for         5041       377.C1.sp6:141918       ATPases       4       655       5933       for         3404       176.F10.sp6:134581       Bcl-2       69       356       16419       for         4036       367.F5.sp6:141182       bromodomain       40       210       8810       for         4489       369.D3.sp6:141543       bromodomain       63       230       10270       for         3408       172.E1.sp6:133925       BZIP       146       298       4066       for         3951       393.G5.sp6:148976       BZIP       116       304       5931       for         4850       172.E9.sp6:133993       BZIP       91       260       4366       for         3895       395.G6.sp6:149361       cyclin       11       281       6930       for         4536       395.G8.sp6:149363       cyclin       12       279       5950 <td< td=""><td></td><td>•</td><td></td><td></td><td></td><td></td><td></td></td<>		•					
4792       184.C6.sp6:135555       ATPases       333       813       5773       for         4847       184.B11.sp6:135548       ATPases       14       498       6140       for         5041       377.C1.sp6:141918       ATPases       4       655       5933       for         3404       176.F10.sp6:134581       Bcl-2       69       356       16419       for         4036       367.F5.sp6:141182       bromodomain       40       210       8810       for         4489       369.D3.sp6:141543       bromodomain       63       230       10270       for         3408       172.E1.sp6:133925       BZIP       146       298       4066       for         395.1       393.G5.sp6:148976       BZIP       116       304       5931       for         4850       172.E9.sp6:133933       BZIP       91       260       4366       for         3618       370.B12.sp6:141721       cyclin       118       324       8980       for         3895       395.G6.sp6:149361       cyclin       11       281       6930       for         4536       395.G8.sp6:149363       cyclin       12       279       5950 <t< td=""><td></td><td><del>-</del></td><td></td><td></td><td></td><td></td><td></td></t<>		<del>-</del>					
4847       184.B11.sp6:135548       ATPases       14       498       6140       for         5041       377.C1.sp6:141918       ATPases       4       655       5933       for         3404       176.F10.sp6:134581       Bcl-2       69       356       16419       for         4036       367.F5.sp6:141182       bromodomain       40       210       8810       for         4489       369.D3.sp6:141543       bromodomain       63       230       10270       for         3408       172.E1.sp6:133925       BZIP       146       298       4066       for         395.1       393.G5.sp6:148976       BZIP       116       304       5931       for         4850       172.E9.sp6:133933       BZIP       91       260       4366       for         3818       370.B12.sp6:141721       cyclin       118       324       8980       for         3895       395.G6.sp6:149361       cyclin       11       281       6930       for         4536       395.G8.sp6:149363       cyclin       12       279       5950       for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479		-					
5041         377.C1.sp6:141918         ATPases         4         655         5933         for           3404         176.F10.sp6:134581         Bcl-2         69         356         16419         for           4036         367.F5.sp6:141182         bromodomain         40         210         8810         for           4489         369.D3.sp6:141543         bromodomain         63         230         10270         for           3408         172.E1.sp6:133925         BZIP         146         298         4066         for           3951         393.G5.sp6:148976         BZIP         116         304         5931         for           4850         172.E9.sp6:133933         BZIP         91         260         4366         for           3618         370.B12.sp6:149361         cyclin         118         324         8980         for           4536         395.G6.sp6:149361         cyclin         11         281         6930         for           4536         395.G8.sp6:149363         cyclin         12         279         5950         for           4455         99.F5.sp6:131294         Cys-protease         72         348         18479         for							
3404       176.F10.sp6:134581       Bcl-2       69       356       16419       for         4036       367.F5.sp6:141182       bromodomain       40       210       8810       for         4489       369.D3.sp6:141543       bromodomain       63       230       10270       for         3408       172.E1.sp6:133925       BZIP       146       298       4066       for         3951       393.G5.sp6:148976       BZIP       116       304       5931       for         4850       172.E9.sp6:133933       BZIP       91       260       4366       for         3618       370.B12.sp6:141721       cyclin       118       324       8980       for         3895       395.G6.sp6:149361       cyclin       11       281       6930       for         4536       395.G8.sp6:149363       cyclin       12       279       5950       for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479       for         4688       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4801       177.E4.sp6:134755       Cys-protease       48       326       1999							
4036       367.F5.sp6:141182       bromodomain       40       210       8810       for         4489       369.D3.sp6:141543       bromodomain       63       230       10270       for         3408       172.E1.sp6:133925       BZIP       146       298       4066       for         3951       393.G5.sp6:148976       BZIP       116       304       5931       for         4850       172.E9.sp6:133933       BZIP       91       260       4366       for         3618       370.B12.sp6:141721       cyclin       118       324       8980       for         3895       395.G6.sp6:149361       cyclin       12       279       5950       for         4536       395.G8.sp6:149363       cyclin       12       279       5950       for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479       for         4684       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4688       180.D1.sp6:135951       Cys-protease       48       326       19999       for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702 <t< td=""><td></td><td>-</td><td></td><td></td><td></td><td></td><td></td></t<>		-					
4489       369.D3.sp6:141543       bromodomain       63       230       10270       for         3408       172.E1.sp6:133925       BZIP       146       298       4066       for         3951       393.G5.sp6:148976       BZIP       116       304       5931       for         4850       172.E9.sp6:133933       BZIP       91       260       4366       for         3618       370.B12.sp6:141721       cyclin       118       324       8980       for         3895       395.G6.sp6:149361       cyclin       11       281       6930       for         4536       395.G8.sp6:149363       cyclin       12       279       5950       for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479       for         4684       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4688       180.D1.sp6:135951       Cys-protease       48       326       19999       for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290       rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828		=					
3408       172.E1.sp6:133925       BZIP       146       298       4066       for         3951       393.G5.sp6:148976       BZIP       116       304       5931       for         4850       172.E9.sp6:133933       BZIP       91       260       4366       for         3618       370.B12.sp6:141721       cyclin       118       324       8980       for         3895       395.G6.sp6:149361       cyclin       11       281       6930       for         4536       395.G8.sp6:149363       cyclin       12       279       5950       for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479       for         4684       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4688       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4801       177.E4.sp6:134755       Cys-protease       48       326       19999       for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290       rev         4821       377.C8.sp6:14179       EFhand       79       153       378		-					
3951       393.G5.sp6:148976       BZIP       116       304       5931       for         4850       172.E9.sp6:133933       BZIP       91       260       4366       for         3618       370.B12.sp6:141721       cyclin       118       324       8980       for         3895       395.G6.sp6:149361       cyclin       12       279       5950       for         4536       395.G8.sp6:149363       cyclin       12       279       5950       for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479       for         4684       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4688       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4801       177.E4.sp6:134755       Cys-protease       48       326       19999       for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290       rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867       rev         5083       216.A1.sp6:134779       EFhand       79       153							
4850       172.E9.sp6:133933       BZIP       91       260       4366       for         3618       370.B12.sp6:141721       cyclin       118       324       8980       for         3895       395.G6.sp6:149361       cyclin       11       281       6930       for         4536       395.G8.sp6:149363       cyclin       12       279       5950       for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479       for         4684       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4688       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4801       177.E4.sp6:134755       Cys-protease       48       326       19999       for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290       rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867       rev         5083       216.A1.sp6:134779       EFhand       79       153       3780       for         2893       185.A1.sp6:135718       EFhand       287       358		<del>-</del>					
3618       370.B12.sp6:141721       cyclin       118       324       8980 for         3895       395.G6.sp6:149361       cyclin       11       281       6930 for         4536       395.G8.sp6:149363       cyclin       12       279       5950 for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479 for         4684       180.D1.sp6:135951       Cys-protease       38       992       10103 rev         4688       180.D1.sp6:135951       Cys-protease       38       992       10103 rev         4801       177.E4.sp6:134755       Cys-protease       48       326       19999 for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290 rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867 rev         5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532 for         2734       177.G4.sp6:134779       EFhand       79       153       3780 for         2893       185.A1.sp6:135718       EFhand       287       358       2580 rev         377.A5.sp6:141136       EFhand       477       563       3010 for		<del>-</del>					for
3895       395.G6.sp6:149361       cyclin       11       281       6930       for         4536       395.G8.sp6:149363       cyclin       12       279       5950       for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479       for         4684       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4688       180.D1.sp6:135951       Cys-protease       38       992       10103       rev         4801       177.E4.sp6:134755       Cys-protease       48       326       19999       for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290       rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867       rev         5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532       for         2734       177.G4.sp6:134779       EFhand       287       358       2580       rev         3775       377.A5.sp6:141898       EFhand       225       272       2500       rev         4152       218.B10.sp6:138803       EFhand       40 <t< td=""><td></td><td><u>-</u></td><td>cyclin</td><td></td><td></td><td></td><td>for</td></t<>		<u>-</u>	cyclin				for
4536       395.G8.sp6:149363       cyclin       12       279       5950 for         4455       99.F5.sp6:131294       Cys-protease       72       348       18479 for         4684       180.D1.sp6:135951       Cys-protease       38       992       10103 rev         4688       180.D1.sp6:135951       Cys-protease       38       992       10103 rev         4801       177.E4.sp6:134755       Cys-protease       48       326       19999 for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290 rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867 rev         5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532 for         2734       177.G4.sp6:134779       EFhand       79       153       3780 for         2893       185.A1.sp6:135718       EFhand       287       358       2580 rev         3775       377.A5.sp6:141898       EFhand       477       563       3010 for         4056       367.B7.sp6:141136       EFhand       40       114       2640 rev         4153       218.B10.sp6:138803       EFhand       40       114		<u>-</u>	•				for
4455       99.F5.sp6:131294       Cys-protease       72       348       18479 for         4684       180.D1.sp6:135951       Cys-protease       38       992       10103 rev         4688       180.D1.sp6:135951       Cys-protease       38       992       10103 rev         4801       177.E4.sp6:134755       Cys-protease       48       326       19999 for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290 rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867 rev         5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532 for         2734       177.G4.sp6:134779       EFhand       79       153       3780 for         2893       185.A1.sp6:135718       EFhand       287       358       2580 rev         3775       377.A5.sp6:141898       EFhand       477       563       3010 for         4056       367.B7.sp6:141136       EFhand       40       114       2640 rev         4153       218.B10.sp6:138803       EFhand       40       114       2640 rev         4154       218.C10.sp6:138815       EFhand       39       113 <td></td> <td>-</td> <td>•</td> <td>12</td> <td>279</td> <td>5950</td> <td>for</td>		-	•	12	279	5950	for
4684       180.D1.sp6:135951       Cys-protease       38       992       10103 rev         4688       180.D1.sp6:135951       Cys-protease       38       992       10103 rev         4801       177.E4.sp6:134755       Cys-protease       48       326       19999 for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290 rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867 rev         5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532 for         2734       177.G4.sp6:134779       EFhand       79       153       3780 for         2893       185.A1.sp6:135718       EFhand       287       358       2580 rev         3775       377.A5.sp6:141898       EFhand       477       563       3010 for         4056       367.B7.sp6:141136       EFhand       225       272       2500 rev         4152       218.B10.sp6:138803       EFhand       40       114       2640 rev         4153       218.C10.sp6:138815       EFhand       40       114       2640 rev         4154       218.C10.sp6:138815       EFhand       39       113	4455	_	Cys-protease	72	348	18479	for
4688       180.D1.sp6:135951       Cys-protease       38       992       10103 rev         4801       177.E4.sp6:134755       Cys-protease       48       326       19999 for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290 rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867 rev         5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532 for         2734       177.G4.sp6:134779       EFhand       79       153       3780 for         2893       185.A1.sp6:135718       EFhand       287       358       2580 rev         3775       377.A5.sp6:141898       EFhand       477       563       3010 for         4056       367.B7.sp6:141136       EFhand       225       272       2500 rev         4152       218.B10.sp6:138803       EFhand       40       114       2640 rev         4153       218.C10.sp6:138815       EFhand       39       113       2640 rev	4684	-	· -		992		
4801       177.E4.sp6:134755       Cys-protease       48       326       19999 for         4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290 rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867 rev         5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532 for         2734       177.G4.sp6:134779       EFhand       79       153       3780 for         2893       185.A1.sp6:135718       EFhand       287       358       2580 rev         3775       377.A5.sp6:141898       EFhand       477       563       3010 for         4056       367.B7.sp6:141136       EFhand       225       272       2500 rev         4152       218.B10.sp6:138803       EFhand       40       114       2640 rev         4153       218.B10.sp6:138803       EFhand       40       114       2640 rev         4154       218.C10.sp6:138815       EFhand       39       113       2640 rev	4688	-	· •	38	992		rev
4659       100.G6.sp6:131499       DAG_PE_bind       605       702       6290       rev         4821       377.C8.sp6:141925       Dead_box_helic       172       828       7867       rev         5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532       for         2734       177.G4.sp6:134779       EFhand       79       153       3780       for         2893       185.A1.sp6:135718       EFhand       287       358       2580       rev         3775       377.A5.sp6:141898       EFhand       477       563       3010       for         4056       367.B7.sp6:141136       EFhand       225       272       2500       rev         4152       218.B10.sp6:138803       EFhand       40       114       2640       rev         4153       218.B10.sp6:138803       EFhand       40       114       2640       rev         4154       218.C10.sp6:138815       EFhand       39       113       2640       rev	4801	-		48	326	19999	for
5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532 for         2734       177.G4.sp6:134779       EFhand       79       153       3780 for         2893       185.A1.sp6:135718       EFhand       287       358       2580 rev         3775       377.A5.sp6:141898       EFhand       477       563       3010 for         4056       367.B7.sp6:141136       EFhand       225       272       2500 rev         4152       218.B10.sp6:138803       EFhand       40       114       2640 rev         4153       218.B10.sp6:138803       EFhand       40       114       2640 rev         4154       218.C10.sp6:138815       EFhand       39       113       2640 rev	4659	100.G6.sp6:131499		605	702	6290	rev
5083       216.A1.sp6:139166       Dead_box_helic       44       589       26532 for         2734       177.G4.sp6:134779       EFhand       79       153       3780 for         2893       185.A1.sp6:135718       EFhand       287       358       2580 rev         3775       377.A5.sp6:141898       EFhand       477       563       3010 for         4056       367.B7.sp6:141136       EFhand       225       272       2500 rev         4152       218.B10.sp6:138803       EFhand       40       114       2640 rev         4153       218.B10.sp6:138803       EFhand       40       114       2640 rev         4154       218.C10.sp6:138815       EFhand       39       113       2640 rev	4821	377.C8.sp6:141925	Dead box helic	172	828	7867	rev
2734       177.G4.sp6:134779       EFhand       79       153       3780 for         2893       185.A1.sp6:135718       EFhand       287       358       2580 rev         3775       377.A5.sp6:141898       EFhand       477       563       3010 for         4056       367.B7.sp6:141136       EFhand       225       272       2500 rev         4152       218.B10.sp6:138803       EFhand       40       114       2640 rev         4153       218.B10.sp6:138803       EFhand       40       114       2640 rev         4154       218.C10.sp6:138815       EFhand       39       113       2640 rev	5083	<u>-</u>		44	589	26532	for
2893       185.A1.sp6:135718       EFhand       287       358       2580       rev         3775       377.A5.sp6:141898       EFhand       477       563       3010       for         4056       367.B7.sp6:141136       EFhand       225       272       2500       rev         4152       218.B10.sp6:138803       EFhand       40       114       2640       rev         4153       218.B10.sp6:138803       EFhand       40       114       2640       rev         4154       218.C10.sp6:138815       EFhand       39       113       2640       rev	2734	<del>-</del>	EFhand	79	153	3780	for
4056       367.B7.sp6:141136       EFhand       225       272       2500       rev         4152       218.B10.sp6:138803       EFhand       40       114       2640       rev         4153       218.B10.sp6:138803       EFhand       40       114       2640       rev         4154       218.C10.sp6:138815       EFhand       39       113       2640       rev	2893	185.A1.sp6:135718	EFhand	287	358	2580	rev
4056       367.B7.sp6:141136       EFhand       225       272       2500       rev         4152       218.B10.sp6:138803       EFhand       40       114       2640       rev         4153       218.B10.sp6:138803       EFhand       40       114       2640       rev         4154       218.C10.sp6:138815       EFhand       39       113       2640       rev			EFhand	477	563	3010	for
4152       218.B10.sp6:138803       EFhand       40       114       2640       rev         4153       218.B10.sp6:138803       EFhand       40       114       2640       rev         4154       218.C10.sp6:138815       EFhand       39       113       2640       rev		•					
4153 218.B10.sp6:138803 EFhand 40 114 2640 rev 4154 218.C10.sp6:138815 EFhand 39 113 2640 rev		-					
4154 218.C10.sp6:138815 EFhand 39 113 2640 rev		<u>-</u>					
		-					
		393.H12.sp6:148995					
4943 219.A9.sp6:138982 EFhand 685 750 2550 rev	4943		EFhand	685	750	2550	rev

SEQ ID NO:	Validation Sequence	Biological Activity (Profile)	Start	Stop	Score	Direction
2849	218.B5.sp6:138798	Ets Nterm	340	531	10400	for
2728	180.A2.sp6:135916	FNtypeII	291	423	6400	rev
3018	216.C1.sp6:139190	FNtypeII	501	634	6460	for
4496	218.G1.sp6:138854	FNtypeII	20	141	6180	rev
4914	393.H8.sp6:148991	FNtypeII	448	576	6110	for
2504	181.C3.sp6:135298	G-alpha	66	715	8084	rev
3290	176.C6.sp6:134541	G-alpha	62	690	9062	for
4288	121.B4.sp6:131896	G-alpha	46	447	21415	for
4444	217.D12.sp6:139405	G-alpha	15	702	40404	for
4562	404.B7.sp6:162874	G-alpha	120	682	8424	for
2503	180.A11.sp6:135925	helicase C	165	479	4494	for
4469	369.C4.sp6:141532	helicase_C	559	756	3732	rev
5020	185.D12.sp6:135765	helicase C	381	534	5000	for
4241	396.H8.sp6:149567	homeobox	80	230	5170	for
2550	180.E5.sp6:135967	mkk	342	612	5791	for
3407	172.F1.sp6:133937	mkk	94	669	5688	rev
3451	123.A2.sp6:132266	mkk	26	378	7889	for
3600	394.B3.sp6:149106	mkk	32	782	9544	for
3646	370.H4.sp6:141785	mkk	18	307	9394	for
3680	369.G11.sp6:141587	mkk	182	725	5375	for
4175	219.H10.sp6:139067	mkk	280	723	15454	for
4205	368.B5.sp6:141327	mkk	249	725	5502	for
4278	181.C9.sp6:135304	mkk	168	880	5551	rev
4322	121.F6.sp6:131946	mkk	111	730	5399	for
4777	177.E2.sp6:134753	mkk	288	636	5720	rev
4482	100.D2.sp6:131459	PDEase	849	1195	5945	for
2578	181.H11.sp6:135366	protkinase	116	710	5531	for
2712	177.G7.sp6:134782	protkinase	6	511	5445	for
2835	218.C1.sp6:138806	protkinase	127	747	5492	for
2843	218.E1.sp6:138830	protkinase	64	726	5592	rev
2971	217.F4.sp6:139421	protkinase	83	702	5818	rev
3009	217.A4.sp6:139361	protkinase	57	682	5395	rev
3084	121.E2.sp6:131930	protkinase	69	658	5593	rev
3226	100.D8.sp6:131465	protkinase	174	620	5453	for
3274	100.C3.sp6:131448	protkinase	228	736	5616	for
3356	172.B5.sp6:133893	protkinase	148	715	5381	for
3377	172.B6.sp6:133894	protkinase	119	775	5616	for
3451	123.A2.sp6:132266	protkinase	24	384	9797	for
3600	394.B3.sp6:149106	protkinase	357	780	11395	for
3635	377.G11.sp6:141976	protkinase	117	739	5992	for
3646	370.H4.sp6:141785	protkinase	24	275	8338	for
3665	370.F2.sp6:141759	protkinase	33	800	5658	for
3669	369.B10.sp6:141526	protkinase	1	482	5504	rev
3700	369.D2.sp6:141542	protkinase	28	661	5428	for
3710	369.G6.sp6:141582	protkinase	71	631	5751	for
3791	396.C11.sp6:149510	protkinase	27	709	5793	rev
3905	393.H7.sp6:148990	protkinase	88	680	5470	rev
3919	393.D10.sp6:148945	protkinase	72	594	5617	for
4044	367.G4.sp6:141193	protkinase	30	699	5439	for
4072	368.B2.sp6:141324	protkinase	44	800	5556	for

	Validation Sequence	Biological	Start	Stop	Score	Direction
NO:	210 D11 6.120020	Activity (Profile)	20	701	(422	<b>C</b>
4117	218.D11.sp6:138828	protkinase	38	781	6423	for
4175	219.H10.sp6:139067	protkinase	277	717	15720	for
4373	216.E5.sp6:139218	protkinase	115	710	5537	for
4569	100.C10.sp6:131455	protkinase	56	783	5556	rev
4755	370.F4.sp6:141761	protkinase	39	803	5635	for
4760	370.F3.sp6:141760	protkinase	188	775	5771	for
4807	184.H3.sp6:135612	protkinase	23	699	5515	for
5059	180.B5.sp6:135931	protkinase	182	671	5718	rev
5102	393.F4.sp6:148963	protkinase	28	650	5345	for
3671	369.D10.sp6:141550	ras	12	332	9802	for
3936	393.A3.sp6:148902	Thioredox	0	263	5887	rev
3927	393.F11.sp6:148970	TNFR c6	151	261	6445	for
2956	184.E10.sp6:135583	transmembrane4	19	483	8339	rev
2981	217.E6.sp6:139411	transmembrane4	83	728	8417	for
3836	396.C9.sp6:149508	transmembrane4	300	924	9444	rev
4038	367.A6.sp6:141123	transmembrane4	32	495	8407	rev
4364	123.A1.sp6:132265	transmembrane4	1289	1548	8114	rev
4406	122.C1.sp6:132097	transmembrane4	6	535	8122	for
4431	122.E4.sp6:132124	transmembrane4	10	530	8829	for
4441	99.F7.sp6:131296	transmembrane4	613	1253	9443	rev
4442	99.F7.sp6:131296	transmembrane4	613	1253	9443	
4653	100.D7.sp6:131464	transmembrane4	335	1207		rev
4654	100.D7.sp6:131464	transmembrane4	335		8255	rev
4710	367.H9.sp6:141210	transmembrane4	398	1207	8255	rev
4944	180.H7.sp6:136005			1130	8352	rev
3381	<del>-</del>	transmembrane4	356	983	8356	rev
	176.D9.sp6:134556	trypsin	164	764	9670	rev
4684	180.D1.sp6:135951	trypsin	371	1229	10479	rev
4688	180.D1.sp6:135951	trypsin	371	1229	10479	rev
2754	177.H6.sp6:134793	WD_domain	345	437	6510	for
3046	121.B8.sp6:131900	WD_domain	98	193	6400	for
3227	100.B10.sp6:131443	WD_domain	544	642	6590	for
4243	121.A8.sp6:131888	WD_domain	93	188	6400	for
5046	185.F10.sp6:135787	WD_domain	382	480	5880	for
3129	121.E12.sp6:131940	Wnt_dev_sign	101	821	12160	rev
3173	99.G6.sp6:131307	Wnt_dev_sign	49	880	12334	rev
3390	176.C9.sp6:134544	Wnt_dev_sign	249	854	11038	rev
3391	176.C9.sp6:134544	Wnt_dev_sign	249	854	11038	rev
3656	370.G6.sp6:141775	Wnt_dev_sign	211	785	11490	rev
3836	396.C9.sp6:149508	Wnt_dev_sign	282	1017	12318	rev
4253	396.F9.sp6:149544	Wnt_dev_sign	482	1298	11217	rev
4330	122.A2.sp6:132074	Wnt_dev_sign	94	933	12383	rev
4359	123.B2.sp6:132278	Wnt_dev_sign	538	1435	11785	for
4364	123.A1.sp6:132265	Wnt_dev_sign	760	1544	12660	rev
4375	122.G10.sp6:132154	Wnt_dev_sign	29	884	11603	rev
4385	122.A2.sp6:132074		94	933	12383	rev
4409	121.F12.sp6:131952	Wnt_dev_sign	9	734	11167	rev
4441	99.F7.sp6:131296	Wnt_dev_sign	560	1399	13749	rev
4442	99.F7.sp6:131296	Wnt_dev_sign	560	1399	13749	rev
4535	395.F10.sp6:149353	Wnt_dev_sign	100	907	11535	rev
4586	123.A4.sp6:132268	Wnt_dev_sign	80	1122	11249	rev

SEQ ID	Validation Sequence	Biological	Start	Stop	Score	Direction
NO:		Activity (Profile)				
4605	404.D5.sp6:162896	Wnt_dev_sign	31	816	11304	rev
4653	100.D7.sp6:131464	Wnt_dev_sign	467	1314	11882	rev
4654	100.D7.sp6:131464	Wnt_dev_sign	467	1314	11882	rev
4665	177.B11.sp6:134726	Wnt_dev_sign	137	1266	12708	rev
4668	177.B11.sp6:134726	Wnt_dev_sign	137	1266	12708	rev
4682	177.B11.sp6:134726	Wnt_dev_sign	137	1266	12708	rev
4710	367.H9.sp6:141210	Wnt_dev_sign	692	1481	12886	rev
4718	185.D9.sp6:135762	Wnt_dev_sign	129	890	11145	rev
4724	377.D2.sp6:141931	Wnt_dev_sign	400	1227	11044	rev
4733	185.D9.sp6:135762	Wnt_dev_sign	129	890	11145	rev
4856	367.H11.sp6:141212	Wnt_dev_sign	295	1669	13366	rev
4866	377.D4.sp6:141933	Wnt_dev_sign	549	1380	14522	rev
4925	219.B12.sp6:138997	Wnt_dev_sign	312	1214	13188	rev
4959	219.B12.sp6:138997	Wnt_dev_sign	312	1214	13188	rev
3409	172.D1.sp6:133913	Y_phosphatase	476	804	6932	for
3418	123.F9.sp6:132333	Y_phosphatase	28	439	6096	rev
3419	123.F9.sp6:132333	Y_phosphatase	28	439	6096	rev
3657	370.H6.sp6:141787	Y_phosphatase	148	554	6481	for
3804	404.B10.sp6:162877	Y_phosphatase	104	466	6446	rev
3806	404.D10.sp6:162901	Y_phosphatase	9	614	6516	for
3974	395.F2.sp6:149345	Y_phosphatase	164	645	6093	rev
4238	121.E9.sp6:131937	Y_phosphatase	240	777	6147	rev
4263	216.F10.sp6:139235	Y_phosphatase	21	<b>50</b> 4	6342	for
4343	122.E9.sp6:132129	Y_phosphatase	381	807	6036	rev
4363	123.B1.sp6:132277	Y_phosphatase	61	510	6229	rev
4434	219.F4.sp6:139037	Y_phosphatase	2	261	10353	for
4473	369.B4.sp6:141520	Y_phosphatase	231	768	6110	rev
4629	404.E11.sp6:162914	Y_phosphatase	580	920	6005	rev
5094	217.A3.sp6:139360	Y_phosphatase	263	622	6222	rev
2738	177.A6.sp6:134709	Zincfing_C2H2	65	127	4380	for
2760	177.A6.sp6:134709	Zincfing_C2H2	65	127	4380	for
2832	218.B2.sp6:138795	Zincfing_C2H2	94	156	4940	for
3736	377.H8.sp6:141985	Zincfing_C2H2	495	557	4850	for
3762	377.G2.sp6:141967	Zincfing_C2H2	52	114	4380	for
3763	377.G2.sp6:141967	Zincfing_C2H2	52	114	4380	for
4794	377.G4.sp6:141969	Zincfing_C2H2	247	308	3930	for
5090	185.C4.sp6:135745	Zincfing_C2H2	238	300	4540	for
3774	377.E4.sp6:141945	Zincfing_C3HC4	128	244	9335	for
4477	181.E3.sp6:135322	Zincfing_C3HC4	321	445	8221	for

Table 19. Polynucleotides Specifically Expressed in Colon											
SEQ	Sequence Name	cluster	lib 1	lib 2	lib 15			lib 18		lib 20	
ID NO:			ciones	ciones	clones	clones	clones	ciones	clones	clones	
3	RTA00000197AF.e.24.1	39250	2	0	0	0	0	0	0	0	
7	RTA00000197AR.e.12.1	22095	3	0	0	0	0	0	0	0	
16	RTA00000196AF.e.16.1	39252	2	0	0	0	0	0	0	0	
18	RTA00000196AF.c.17.1	39602	2	0	0	0	0	0	0	0	
21	RTA00000131A.g.19.2	36535	2	0	0	0	0	0	0	0	
22	RTA00000187AR.o.10.2	8984	4	3	0	0	0	2	0	0	
23	RTA00000198R.b.08.1	22636	3	0	0	0	0	0	0	0	
26	RTA00000200R.g.09.1	22785	3	0	0	0	0	0	0	0	
29	RTA00000200AF.b.19.1	22847	3	0	0	0	0	0	0	0	
31	RTA00000200F.m.15.1	22601	3	0	0	0	1	0	0	0	
37	RTA00000181AF.n.15.2	86128	1	0	0	0	0	0	0	0	
38	RTA00000196R.k.07.1	22443	2	0	0	0	0	0	0	1	
40	RTA00000200AR.e.02.1	36059	2	0	0	0	1	1	1	0	
48	RTA00000177AR.a.23.5	6995	4	2	0	0	0	0	0	0	
49	RTA00000198R.o.05.1	26702	2	0	0	0	0	0	0	0	
50	RTA00000201R.a.02.1	35362	2	0	0	0	0	0	0	0	
61	RTA00000197AF.h.11.1	22264	3	0	0	0	0	0	0	0	
66	RTA00000199F.c.09.2	16824	3	1	0	0	0	0	0	0	
75	RTA00000180AR.h.19.2	84182	1	0	0	0	0	0	0	0	
78	RTA00000199R.f.09.1	22907	3	0	0	0	0	0	0	0	
79	RTA00000199AF.p.4.1	10282	3	3	0	0	0	0 -	0	0	
85	RTA00000200R.o.03.1	22807	3	0	0	0	0	0	0	0	
86	RTA00000189AF.1.22.1	33333	1	1	0	0	0	0	0	0	
87	RTA00000195AF.d.20.1	37574	2	0	0	0	0	0	0	0	
92	RTA00000198AF.j.18.1	22759	3	0	0	0	0	0	0	0	
95	RTA00000180AF.g.3.1	9024	5	2	0	0	0	0	0	0	
102	RTA00000199R.j.08.1	37844	2	0	0	0	0	0	0	0	
103	RTA00000199F.e.10.1	22906	3	0	0	0	0	0	1	0	
105	RTA00000179AF.g.12.3	36390	2	0	0	0	0	0	0	0	
108	RTA00000183AR.h.23.2	18957	3	0	0	0	0	0	0	0	
109	RTA00000197AF.d.12.1	39546	2	0	0	0	0	0	0	0	
116	RTA00000181AR.k.24.3	7005	8	2	0	0	0	0	0	0	
119	RTA00000181AR.k.24.2	7005	8	2	0	0	0	0	0	0	
124	RTA00000199AR.m.06.1	19122	3	0	0	0	0	0	0	0	
129	RTA00000134A.d.10.1	18957	3	0	0	0	0	0	0	0	
137	RTA00000181AF.m.4.3	13238	4	1	0	0	0	0	0	0	
141	RTA00000196AF.c.6.1	23148	3	0	0	0	0	0	0	0	
142	RTA00000198AF.k.19.1	75879	1	0	0	0	0	0	0	0	
				602							

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones		lib 16 clones				lib 20 clones
143	RTA00000199R.h.09.1	76020	1	0	0	0	0	0	0	0
144	RTA00000198AF.o.18.1	13018	4	0	0	0	1	0	0	0
148	RTA00000199F.h.17.2	36254	2	0	0	0	0	0	0	0
149	RTA00000181AR.h.06.3	87226	1	0	0	0	0	0	0	0
166	RTA00000198AF.f.21.1	22676	3	0	0	0	0	0	0	0
173	RTA00000200AR.b.07.1	17125	4	0	0	0	0	0	0	0
178	RTA00000200F.o.03.1	22807	3	0	0	0	0	0	0	0
180	RTA00000199AF.j.12.1	22461	3	0	0	0	0	0	0	0
185	RTA00000195AF.d.4.1	22766	3	0	0	0	0	0	0	0
194	RTA00000200R.k.01.1	40049	2	0	0	0	0	0	0	0
195	RTA00000198AF.c.10.1	77149	1	0	0	0	0	0	0	0
198	RTA00000197AR.e.07.1	86969	1	0	0	0	0	0	0	0
199	RTA00000199R.c.09.1	16824	3	1	0	0	0	0	0	0
206	RTA00000181AF.o.04.2	22205	3	0	0	0	0	0	0	0
207	RTA00000199AF.l.19.1	22460	3	0	0	0	0	0	0	0
208	RTA00000198AF.h.22.1	22366	2	1	0	0	0	0	0	0
211	RTA00000199AF.m.15.1	10101	3	0	0	0	0	0	0	0
212	RTA00000197AF.j.9.1	13236	4	1	0	0	0	0	0	0
230	RTA00000185AR.b.18.1	12171	3	2	0	0	0	0	0	0
235	RTA00000201AF.a.02.1	35362	2	0	0	0	0	0	0	0
236	RTA00000183AR.h.23.1	18957	3	0	0	0	0	0	0	0
238	RTA00000187AR.k.12.1	78415	1	0	0	0	0	0 .	0	0
242	RTA00000198AF.m.17.1	77992	1	0	0	0	0	0	0	0
243	RTA00000181AF.m.15.3	12081	4	0	0	0	0	0	0	0
248	RTA00000198R.c.14.1	39814	2	0	0	0	0	0	0	0
249	RTA00000200R.o.03.2	22807	3	0	0	0	0	0	0	0
251	RTA00000192AF.n.13.1	8210	2	6	0	0	0	0	0	0
256	RTA00000184AR.e.15.1	16347	4	0	0	0	0	0	0	0
260	RTA00000198R.m.17.1	77992	1	0	0	0	0	0	0	0
270	RTA00000178R.I.08.1	39648	2	0	0	0	0	0	0	0
278	RTA00000198AF.p.16.1	71877	1	0	0	0	0	0	0	0
280	RTA00000193AF.b.18.1	7542	8	0	0	2	1	0	1	0
284	RTA00000199F.d.10.2	22049	3	0	0	0	0	0	0	0
287	RTA00000200AF.b.07.1	17125	4	0	0	0	0	0	0	0
288	RTA00000181AR.i.06.3	19119	3	0	0	0	0	0	0	0
289	RTA00000196F.k.07.1	22443	2	0	0	0	0	0	0	1
294	RTA00000198AF.k.23.1	8995	2	5	0	0	0	0	0	0
296	RTA00000196AF.f.20.1	22774	3	0	0	0	0	0	0	0
300	RTA00000195AF.c.12.1	37582	2	0	0	0	0	0	0	0

SEQ ID NO:	•	cluster		lib 2 clones	lib 15	lib 16 clones	lib 17 clones		lib 19 clones	lib 20 clones
302	RTA00000186AF.d.1.2	40044	2	0	0	1	0	0	0	0
307	RTA00000200F.n.05.2	18989	3	0	0	0	0	0	0	0
308	RTA00000178AF.j.20.1	15066	4	0	0	0	0	0	0	0
310	RTA00000188AF.m.08.1	22155	3	0	0	0	0	0	0	0
315	RTA00000199R.d.23.1	37477	2	0	0	0	0	0	0	0
319	RTA00000200F.n.05.1	18989	3	0	0	0	0	0	0	0
320	RTA00000196AF.m.13.1	16290	4	0	0	0	0	0	0	0
325	RTA00000182AF.d.18.4	37435	2	0	0	0	0	0	0	0
328	RTA00000200AF.g.09.1	22785	3	0	0	0	0	0	0	0
330	RTA00000177AR.m.17.4	14391	3	1	0	0	0	0	0	0
331	RTA00000197AR.c.20.1	16282	4	0	0	0	0	0	0	0
337	RTA00000177AR.m.17.3	14391	3	1	0	0	0	0	0	0
342	RTA00000196AF.d.10.1	22256	3	0	0	0	0	0	0	0
343	RTA00000201F.a.18.1	16837	2	2	0	0	0	0	0	0
344	RTA00000198AF.o.02.1	68756	1	0	0	0	0	0	0	0
345	RTA00000187AF.h.21.1	39171	2	0	0	0	0	0	0	0
347	RTA00000199F.b.03.2	38340	2	0	0	0	0	0	0	0
358	RTA00000198AF.g.7.1	13386	3	2	0	0	0	-	0	0
362	RTA00000197AR.c.24.1	82498	1	0	0	0	_		_	0
371	RTA00000197F.e.7.1	86969	1	0	0	0	0			0
378	RTA00000181AF.k.24.3	7005	8	2	0	0	0			0
382	RTA00000200AF.j.6.1	22902	3	0	0	0	0			0
384	RTA00000196AF.h.17.1	39215	2	0	0	0	0			0
392	RTA00000185AF.b.11.2	9024	5	2	0	0	0	0	0	0
397	RTA00000198AF.b.22.1	38956	2	0	0	0	0	0 (	0	0
399	RTA00000186AF.m.15.2	40122	2	0	0	0 (	0 (	0 (	0 (	0
406	RTA00000199F.f.09.2	22907	3	0	0	0 (	0 (	0 (	) (	0
408	RTA00000183AR.1.15.1	39383	2	0	0	0 (	) (	) (	) (	0
413	RTA00000200F.a.12.1	16751	4	0	0	0 (	) (	) (	) (	)
416	RTA00000199F.a.5.1	22134	3	0 (	0	0 (	) (	) (	) (	)
418	RTA00000187AR.k.01.1	78356	1	0 (	0 (	) (	) (	) (	) (	)
424	RTA00000187AR.j.24.1	78356	1	0 (	0 (	) (	) (	) (	) (	)
426	RTA00000199AF.o.19.1	36927	2	0 (	0 (	) (	) (	0	0	)
429	RTA00000196F.i.19.1	39498	2 (	) (	) (	) (	0	0	0	)
430	RTA00000198R.k.23.1	8995	2 :	5 (	) (	) 0	0	0	0	)
432	RTA00000198AF.o.05.1		2 (	) (	) (	0	0	0	0	•
433	RTA00000198R.j.18.1		3 (	) (	) (	0	0	0	0	
435	RTA00000182AR.c.22.1		3 (	) (	) (	0	0	0	0	
438	RTA00000180AR.g.03.4	9024	5 2	2 0	) (	0	0	0	0	

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones		lib 17 clones	lib 18 clones	lib 19 clones	lib 20 clones
451	RTA00000200AF.b.20.1	40403	2	0	0	0	0	0	0	0
455	RTA00000198AF.d.12.1	21142	2	1	0	0	0	0	0	0
456	RTA00000200AF.b.12.1	22053	3	0	0	0	0	0	0	0
457	RTA00000191AR.1.7.2	14391	3	1	0	0	0	0	0	0
461	RTA00000190AF.e.13.1	38961	2	0	0	0	0	0	0	0
462	RTA00000196AF.n.17.1	12477	4	1	0	0	0	0	0	0
467	RTA00000195AF.b.19.1	77678	1	0	0	0	0	0	0	0
475	RTA00000187AR.m.3.3	17055	4	0	0	0	0	0	0	0
476	RTA00000200R.g.15.1	22898	3	0	0	0	0	0	0	0
482	RTA00000187AF.j.7.1	78091	1	0	0	0	0	0	0	0
485	RTA00000196AF.c.14.1	23105	3	0	0	0	0	0	0	0
486	RTA00000190AR.p.22.2	16368	4	0	0	0	0	0	0	0
492	RTA00000198AF.b.8.1	22636	3	0	0	0	0	0	0	0
493	RTA00000177AF.m.17.1	14391	3	1	0	0	0	0	0	0
494	RTA00000200AF.k.1.1	40049	2	0	0	0	0	0	0	0
498	RTA00000190AF.h.12.1	12977	5	0	0	0	0	0	0	0
499	RTA00000199F.b.22.2	17018	4	0	0	0	0	0	0	0
508	RTA00000187AF.i.14.2	19406	2	1	0	0	0	0	0	0
511	RTA00000196AF.g.10.1	12498	3	1	1	0	0	0	0	0
517	RTA00000184AF.e.14.1	16347	4	0	0	0	0	0	0	0
522	RTA00000178AR.h.17.2	23824	2	1	0	0	0	0	0	0
531	RTA00000195F.a.3.1	27179	2	0	0	0	0	0 -	0	0
544	RTA00000196F.j.13.1	23170	3	0	0	0	0	0	0	0
547	RTA00000196AF.g.8.1	39665	2	0	0	0	0	0	0	0
549	RTA00000198AF.c.16.1	26801	2	0	0	0	0	0	0	0
553	RTA00000201F.b.22.1	35728	2	0	0	0	0	0	0	1
559	RTA00000197AF.p.20.1	22795	3	0	0	0	0	0	0	0
563	RTA00000192AR.o.16.2	9061	5	2	0	0	0	0	0	0
565	RTA00000191AF.c.10.1	40422	2	0	0	0	0	0	0	0
568	RTA00000196AF.p.01.2	87143	1	0	0	0	0	0	0	0
578	RTA00000180AF.g.17.1	16653	3	1	0	0	0	0	0	0
583	RTA00000190AR.h.12.2	12977	5	0	0	0	0	0	0	0
585	RTA00000198AF.n.18.1	16715	3	1	0	0	0	0	0	0
586	RTA00000199R.o.11.1	23172	3	0	0	0	0	0	0	0
588	RTA00000191AF.b.4.1	14936	3	0	0	0	0	0	0	0
589	RTA00000192AF.l.1.1	16392	3	0	0	0	0	0	0	0
593	RTA00000196R.c.14.2	23105	3	0	0	0	0	0	0	0
595	RTA00000195R.a.06.1	35265	2	0	1	0	0	0	0	0
602	RTA00000195AF.b.21.1	39055	2	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones		lib 19 clones	lib 20 clones
612	RTA00000197AR.e.22.1	78758	1	0	0	0	0	0	0	0
615	RTA00000197R.p.20.1	22795	3	0	0	0	0	0	0	0
618	RTA00000192AF.a.14.1	6874	6	3	0	0	1	0	0	0
623	RTA00000198R.b.24.1	19047	3	0	0	0	0	0	0	0
627	RTA00000199F.h.15.2	22269	3	0	0	0	0	0	0	0
628	RTA00000198AF.g.16.1	6602	1	1	0	0	0	0	0	0
634	RTA00000192AF.j.6.1	11494	4	0	0	0	0	0	0	0
635	RTA00000181AF.p.7.3	38773	2	0	0	0	0	0	0	0
637	RTA00000200AF.g.15.1	22898	3	0	0	0	0	0	0	0
643	RTA00000184AF.c.9.1	16245	4	0	0	0	0	0	0	0
645	RTA00000177AF.k.9.1	16245	4	0	0	0	0	0	0	0
649	RTA00000190AR.1.19.2	88204	1	0	0	0	0	0	0	0
662	RTA00000201R.a.15.1	57347	1	0	0	0	0	0	0	0
664	RTA00000195R.a.23.1	86432	1	0	0	0	0	0	0	0
670	RTA00000186AF.p.17.3	38383	2	0	0	0	0	0	0	0
674	RTA00000197AR.e.24.1	39250	2	0	0	0	0	0	0	0
683	RTA00000187AR.j.01.1	79028	1	0	0	0	0	0	0	0
686	RTA00000201F.f.07.1	51116	1	0	0	0	0	0	0	0
694	RTA00000201R.c.19.1	22357	2	1	0	0	0	0	0	0
702	RTA00000177AR.b.8.5	17062	3	0	0	0	0	0	0	0
712	RTA00000201F.b.21.1	9071	3	4	0	0	0	0	0	0
717	RTA00000200F.o.10.2	36432	2	0	0	0	0	0 -	0	0
718	RTA00000196F.1.14.2	23144	3	0	0	0	0	0	0	0
725	RTA00000197AF.b.1.1	12134	1	1	0	0	0	0	0	0
733	RTA00000200AF.d.20.1	26600	2	0	0	0	0	0	0	0
743	RTA00000178AF.k.9.1	16342	3	0	0	0	0	0	0	0
748	RTA00000198AF.b.24.1	19047	3	0	0	0	0	0	0	0
757	RTA00000406F.d.16.1	15040	2	2	0	0	0	0	0	0
760	RTA00000408F.o.12.2	78578	1	0	0	0	0	0	0	0
761	RTA00000119A.j.15.1	79623	1	0	0	0	0	0	0	0
762	RTA00000413F.d.12.1	66467	1	0	0	0	0	0	0	0
763	RTA00000423F.i.12.1	9118	4	3	0	0	0	0	0	0
766	RTA00000411F.k.05.1	64777	1	0	0	0	0	0	0	0
769	RTA00000419F.b.09.1	78128	1	0	0	0	0	0	0	0
772	RTA00000411F.m.15.1	78014	1	0	0	0	0	0	0	0
774	RTA00000123A.k.23.1	80313	1	0	0	0	0	0	0	0
777	RTA00000130A.m.15.1	81630	1	0	0	0	0	0	0	0
778	RTA00000411F.k.20.1	64973	1	0	0	0	0	0	0	0
780	RTA00000418F.k.05.1	73021	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones		lib 16 clones			lib 19 clones	lib 20 clones
781	RTA00000423F.h.18.1	37972	2	0	0	0	0	0	0	0
783	RTA00000422F.p.06.2	39282	2	0	0	0	0	0	0	0
784	RTA00000404F.n.16.2	39095	2	0	0	0	0	0	0	0
785	RTA00000411F.m.24.1	77568	1	0	0	0	0	0	0	0
786	RTA00000134A.j.10.1	81383	1	0	0	0	0	0	0	0
787	RTA00000409F.j.02.1	76417	1	0	0	0	0	0	0	0
788	RTA00000403F.j.15.1	23840	2	1	0	0	0	0	0	0
789	RTA00000411F.n.11.1	77276	1	0	0	0	0	0	0	0
790	RTA00000339F.i.13.1	5970	6	4	0	0	0	0	0	0
792	RTA00000406F.o.15.1	37482	2	0	0	0	0	0	0	0
793	RTA00000412F.g.04.2	64457	1	0	0	0	0	0	0	0
795	RTA00000352R.1.06.1	40343	2	0	0	0	0	0	0	0
796	RTA00000419F.b.12.1	63148	1	0	0	0	0	0	0	0
797	RTA00000423F.k.17.2	37512	2	0	0	0	0	0	0	0
799	RTA00000418F.k.14.1	76133	1	0	0	0	0	1	0	0
800	RTA00000409F.l.12.1	26755	1	0	0	0	0	0	0	0
801	RTA00000404F.c.20.1	39088	2	0	0	0	0	0	1	0
802	RTA00000423F.g.09.1	38958	2	0	0	0	0	0	0	0
804	RTA00000406F.d.12.1	38575	2	0	0	0	0	0	0	0
805	RTA00000411F.f.02.1	63386	1	0	0	0	0	0	0	0
806	RTA00000129A.n.21.1	79381	1	0	0	0	0	0	0	0
807	RTA00000409F.m.12.1	73490	1	0	0	0	0	0	-0	0
808	RTA00000410F.c.04.1	74099	1	0	0	0	0	0	0	0
810	RTA00000406F.m.09.1	26891	2	0	0	0	0	0	0	0
811	RTA00000411F.b.06.1	77884	1	0	0	0	0	0	0	0
812	RTA00000409F.1.21.1	73143	1	0	0	0	0	0	0	0
818	RTA00000404F.1.20.2	38638	2	0	0	0	0	0	0	0
819	RTA00000413F.d.18.1	65305	1	0	0	0	0	0	0	0
820	RTA00000404F.p.04.2	39069	2	0	0	0	0	0	0	0
821	RTA00000405F.g.19.2	37150	2	0	0	0	0	0	0	0
822	RTA00000409F.a.22.1	75200	1	0	0	0	0	0	0	0
824	RTA00000405F.o.18.1	11016	4	2	0	0	0	0	0	0
829	RTA00000408F.e.22.2	26930	1	0	0	0	0	0	0	0
831	RTA00000413F.d.16.1	63331	1	0	0	0	0	0	0	0
834	RTA00000419F.g.08.1	66700	1	0	0	0	0	0	0	0
835	RTA00000122A.g.16.1	81366	1	0	0	0	0	0	0	0
836	RTA00000419F.c.16.1	65254	1	0	0	0	0	0	0	0
837	RTA00000411F.b.03.1		1	2	0	0	0	0	0	0
842	RTA00000403F.l.20.1	18267	1	0	0	0	0	0	0	0
				607						

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones		lib 16 clones				lib 20 clones
845	RTA00000411F.a.02.1	78537	1	0	0	0	0	0	0	0
847	RTA00000412F.l.04.1	66372	1	0	0	0	0	0	0	0
849	RTA00000406F.a.23.1	38712	2	0	0	0	0	0	0	0
851	RTA00000120A.n.19.3	80004	1	0	0	0	0	0	0	0
852	RTA00000403F.e.01.1	38965	2	0	0	0	0	0	0	0
853	RTA00000411F.l.03.1	62702	1	0	0	0	0	0	0	0
856	RTA00000121A.m.2.1	81064	1	0	0	0	0	0	0	0
858	RTA00000418F.j.12.1	73316	1	0	0	0	0`	0	0	0
862	RTA00000125A.g.16.1	21497	2	1	0	0	0	0	0	0
863	RTA00000418F.o.18.1	78676	1	0	0	0	0	0	0	0
865	RTA00000408F.k.14.1	73856	1	0	0	0	0	0	0	0
871	RTA00000403F.o.15.1	39140	2	0	0	0	0	0	0	0
872	RTA00000341F.m.13.1	26502	1	0	0	0	0	0	0	0
873	RTA00000408F.h.03.1	78382	1	0	0	0	0	0	0	0
874	RTA00000423F.k.05.1	37472	2	0	0	0	0	0	0	0
876	RTA00000418F.p.19.1	78544	1	0	0	0	0	0	0	0
877	RTA00000420F.f.06.1	64812	1	0	0	0	0	0	0	0
878	RTA00000122A.j.18.1	81317	1	0	0	0	0	0	0	0
879	RTA00000420F.d.05.1	64432	1	0	0	0	0	0	0	0
880	RTA00000403F.m.18.1	39185	2	0	0	0	0	0	0	0
882	RTA00000411F.j.05.1	40709	1	1	0	0	0	0	0	0
883	RTA00000403F.a.04.1	23529	2	1	0	0	0	0 .	0	0
885	RTA00000406F.f.12.1	21895	2	1	0	0	0	0	0	0
886	RTA00000418F.g.22.1	74837	1	0	0	0	0	0	0	0
888	RTA00000404F.1.20.1	38638	2	0	0	0	0	0	0	0
889	RTA00000408F.i.08.2	75811	1	0	0	0	0	0	0	0
890	RTA00000122A.d.5.1	81155	1	0	0	0	0	0	0	0
894	RTA00000419F.b.19.1	65534	1	0	0	0 .	0	0	0	0
896	RTA00000418F.k.19.1	74932	1	0	0	0	0	0	0	0
900	RTA00000419F.g.12.1	66171	1	0	0	0	0	0	0	0
901	RTA00000404F.n.11.2	38001	2	0	0	0	0	0	0	0
904	RTA00000419F.o.24.1	65092	1	0	0	0	0	0	0	0
905	RTA00000419F.k.19.1	75447	1	0	0	0	0	0	0	0
907	RTA00000127A.i.20.1	81418	1	0	0	0	0	0	0	0
908	RTA00000422F.g.22.1	22561	3	0	0	0	0	0	0	0
910	RTA00000413F.h.13.1	65190	1	0	0	0	0	0	0	0
913	RTA00000348R.j.16.1	7005	8	2	0	0	0	0	0	0
916	RTA00000418F.n.22.1	79062	1	0	0	0	0	0	0	0
917	RTA00000406F.1.08.1	39016	2	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones		lib 16 clones			lib 19 clones	lib 20 clones
920	RTA00000409F.j.07.1	75190	1	0	0	0	0	0	0	0
923	RTA00000411F.e.22.1	63638	1	0	0	0	0	0	0	0
924	RTA00000347F.a.17.1	16723	3	1	0	0	0	0	0	0
926	RTA00000404F.n.20.1	26865	2	0	0	0	0	0	0	0
929	RTA00000404F.b.02.1	38984	2	0	0	0	0	0	0	0
931	RTA00000403F.b.10.1	73268	1	0	0	0	0	0	0	0
932	RTA00000406F.i.12.1	39080	2	0	0	0	0	0	0	0
933	RTA00000406F.h.08.1	16228	2	2	0	0	0	0	0	0
934	RTA00000418F.i.19.1	79180	1	0	0	0	0	0	0	0
936	RTA00000412F.h.21.1	64348	1	0	0	0	0	0	0	0
938	RTA00000120A.g.18.1	81255	1	0	0	0	0	0	0	0
940	RTA00000423F.j.05.1	37958	2	0	0	0	0	0	0	0
941	RTA00000132A.k.6.1	81284	1	0	0	0	0	0	0	0
943	RTA00000406F.p.04.1	37458	2	0	0	0	0	0	0	0
944	RTA00000347F.a.13.1	22446	3	0	0	0	0	0	0	0
945	RTA00000419F.p.23.1	64748	1	0	0	0	0	0	0	0
946	RTA00000419F.d.17.1	64353	1	0	0	0	0	0	0	0
949	RTA00000124A.k.5.1	80252	1	0	0	0	0	0	0	0
950	RTA00000404F.h.22.1	18735	2	1	0	0	0	0	1	0
952	RTA00000410F.o.05.1	75262	1	0	0	0	0	0	0	0
953	RTA00000339R.l.14.1	19119	3	0	0	0	0	0	0	0
954	RTA00000403F.m.13.2	39077	2	0	0	0	0	0	0	0
957	RTA00000419F.g.22.1	64515	1	0	0	0	0	0	0	0
958	RTA00000404F.g.21.1	37947	2	0	0	0	0	0	0	0
960	RTA00000138A.n.4.1	21920	2	1	0	0	0	0	0	0
961	RTA00000410F.b.15.1	77100	1	0	0	0	0	0	0	0
963	RTA00000419F.j.23.1	74470	1	0	0	0	0	0	0	0
964	RTA00000411F.j.02.1	65310	1	0	0	0	0	0	0	0
965	RTA00000419F.p.24.1	63477	1	0	0	0	0	0	0	0
966	RTA00000404F.a.19.1	38624	2	0	0	0	0	0	0	0
973	RTA00000346F.e.13.1	74653	1	0	0	0	0	0	0	0
974	RTA00000419F.c.18.1	41394	1	1	0	0	0	0	0	0
978	RTA00000404F.e.22.1	11344	3	3	0	0	0	0	0	0
981	RTA00000125A.k.10.1	81644	1	0	0	0	0	0	0	0
982	RTA00000347F.c.06.1	18846	2	1	0	0	0	0	0	0
983	RTA00000411F.k.19.1	64200	1	0	0	0	0	0	0	0
984	RTA00000345F.i.09.1	27250	2	0	0	0	0	0	0	0
985	RTA00000423F.k.01.1	40426	2	0	0	0	0	0	0	0
986	RTA00000408F.d.06.1	78997	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones		lib 16 clones				lib 20 clones
987	RTA00000128A.b.20.1	79761	1	0	0	0	0	0	0	0
989	RTA00000195AF.d.4.1	22766	3	0	0	0	0	0	0	0
991	RTA00000403F.h.12.1	15205	2	1	0	0	0	0	0	0
992	RTA00000119A.j.22.1	80336	1	0	0	0	0	0	0	0
995	RTA00000126A.n.7.2	79557	1	0	0	1	0	0	0	0
997	RTA00000404F.j.08.1	39066	2	0	0	0	0	0	0	0
998	RTA00000410F.c.14.1	77809	1	0	0	0	0	0	0	0
999	RTA00000120A.g.23.1	81189	1	0	0	0	0	0	0	0
1000	RTA00000195AF.d.20.1	37574	2	0	0	0	0	0	0	0
1002	RTA00000412F.j.17.1	64071	1	0	0	0	0	0	0	0
1004	RTA00000119A.j.10.1	79646	1	0	0	0	0	0	0	0
1010	RTA00000419F.o.16.1	62867	1	0	0	0	0	0	0	0
1012	RTA00000411F.c.17.1	77664	1	0	0	0	0	0	0	0
1013	RTA00000406F.k.15.1	38549	2	0	0	0	0	0	0	0
1014	RTA00000406F.a.02.1	37744	2	0	0	0	0	0	0	0
1016	RTA00000341F.b.06.1	17008	4	0	0	0	0	0	0	0
1017	RTA00000409F.n.14.1	78190	1	0	0	0	0	0	0	0
1019	RTA00000345F.j.08.1	16731	3	1	0	0	0	0	0	0
1021	RTA00000419F.g.15.1	32519	1	1	0	0	0	0	0	0
1022	RTA00000423F.a.19.1	21396	1	2	0	0	0	0	0	0
1024	RTA00000422F.e.08.1	39020	2	0	0	0	0	0	0	0
1025	RTA00000411F.d.15.1	74890	1	0	0	0	0	0 -	0	0
1027	RTA00000411F.l.15.1	66704	1	0	0	0	0	0	0	0
1029	RTA00000405F.e.08.1	37916	2	0	0	0	1	0	0	0
1030	RTA00000353R.j.24.1	23089	3	0	0	0	0	0	0	0
1032	RTA00000418F.o.06.1	75930	1	0	0	0	0	0	0	0
1033	RTA00000404F.c.10.1	23534	2	1	0	0	0	0	0	0
1034	RTA00000418F.i.21.1	78728	1	0	0	0	0	0	0	0
1036	RTA00000411F.I.13.1	43114	1	1	0	0	0	0	0	0
1037	RTA00000407F.a.24.1	37560	2	0	0	0	0	0	0	0
1038	RTA00000346F.n.06.1	12439	4	0	0	0	0	0	0	0
1039	RTA00000412F.1.21.1	65183	1	0	0	0	0	0	0	0
1040	RTA00000413F.i.02.1	65857	1	0	0	0	0	0	0	0
1041	RTA00000404F.i.19.1	38698	2	0	0	0	0	0	0	0
1043	RTA00000403F.a.11.1	73109	1	0	0	0	0	0	0	0
1045	RTA00000411F.k.16.1	64759	1	0	0	0	0	0	1	0
1046	RTA00000405F.c.01.1	19236	2	0	0	0	0	0	0	0
	RTA00000423F.i.18.1	14996	4	0	0	0	0	0	0	0
1050	RTA00000406F.a.07.1	26607	2	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones	lib 18 clones	lib 19 clones	lib 20 clones
1051	RTA00000347F.d.06.1	39122	2	0	0	0	0	0	0	0
1052	RTA00000419F.b.18.1	67034	1	0	0	0	0	0	0	0
1053	RTA00000406F.h.07.1	38003	2	0	0	0	0	0	0	0
1054	RTA00000405F.1.15.1	19575	2	1	0	0	0	0	0	0
1055	RTA00000406F.g.17.1	37979	2	0	0	0	0	0	0	0
1058	RTA00000130A.h.22.1	80933	1	0	0	0	0	0	0	0
1061	RTA00000404F.d.13.1	39036	2	0	0	0	0	0	0	0
1064	RTA00000340F.n.01.1	39081	2	0	0	0	0	0	0	0
1065	RTA00000419F.d.06.1	65496	1	0	0	0	0	0	0	0
1066	RTA00000419F.n.09.1	66070	1	0	0	0	0	0	0	0
1067	RTA00000399F.i.08.1	38927	2	0	0	0	0	0	0	0
1069	RTA00000423F.g.13.1	38028	2	0	0	0	0	0	0	0
1072	RTA00000195AF.b.21.1	39055	2	0	0	0	0	0	0	0
1073	RTA00000403F.h.05.1	39096	2	0	0	0	0	0	0	0
1075	RTA00000422F.p.07.2	39024	2	0	0	1	0	0	0	0
1078	RTA00000421F.n.19.1	16409	3	1	0	0	0	0	0	0
1080	RTA00000345F.k.21.1	40204	2	0	0	0	0	0	0	0
1082	RTA00000405F.a.11.1	39124	2	0	0	0	0	0	0	0
1084	RTA00000413F.e.16.1	63836	1	0	0	0	0	0	0	0
1086	RTA00000404F.o.18.2	39110	2	0	0	0	0	0	0	0
1087	RTA00000409F.i.24.1	76967	1	0	0	0	0	0	0	0
1091	RTA00000340F.n.13.1	17055	4	0	0	0	0	0 -	0	0
1092	RTA00000340F.p.04.1	78533	1	0	0	0	0	0	0	0
1093	RTA00000411F.c.05.1	73368	1	0	0	0	0	0	0	0
1097	RTA00000404F.i.02.1	39015	2	0	0	0	0	0	0	0
1099	RTA00000403F.m.15.2	26901	2	0	0	0	0	0	0	0
1100	RTA00000412F.h.23.2	65118	1	0	0	0	0	0	0	0
1101	RTA00000418F.j.08.1	73382	1	0	0	0	0	0	0	0
1102	RTA00000125A.n.4.1		1	0	0	0	0	0	0	0
1103	RTA00000412F.I.19.1	65825	1	0	0	0	0	0	0	0
1105	RTA00000129A.p.3.1		1	1	0	0	0	0	0	0
1106	RTA00000340F.p.20.1		4	0	0	0	0	0	0	0
1107	RTA00000411F.a.10.1		1	0	0	0	0	0	0	0
1108	RTA00000409F.n.17.1	76725	1	0	0	0	0	0	0	0
1109	RTA00000404F.c.03.2	39198	2	0	0	0	0	0	0	0
1110	RTA00000420F.a.19.1		1	1	0	0,	0	0	0	0
1114	RTA00000420F.d.12.1					-			0	0
1115	RTA00000409F.j.19.1									0
1116	RTA00000422F.d.16.1	39133	2	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones	lib 18 clones	lib 19 clones	lib 20 clones
1117	RTA00000418F.m.16.1	74986	1	0	0	0	0	0	0	0
1118	RTA00000405F.c.11.1	39068	2	0	0	0	0	0	0	0
1119	RTA00000404F.k.22.1	39084	2	0	0	0	0	0	0	0
1120	RTA00000418F.k.07.1	75067	1	0	0	0	0	0	0	0
1121	RTA00000403F.c.10.1	75261	1	0	0	0	0	0	0	0
1124	RTA00000410F.m.05.1	74964	1	0	0	0	0	0	0	0
1125	RTA00000405F.i.20.1	38532	2	0	0	0	0	0	0	0
1127	RTA00000408F.p.24.1	74286	1	0	0	0	0	0	0	0
1128	RTA00000418F.k.18.1	75385	1	0	0	0	0	0	0	0
1129	RTA00000422F.m.04.1	38702	2	0	0	0	0	0	0	0
1133	RTA00000403F.a.07.1	73559	1	0	0	0	0	0	0	0
1135	RTA00000403F.b.19.1	22327	2	1	0	0	0	0	0	0
1136	RTA00000418F.m.23.1	77195	1	0	0	0	0	0	0	0
1138	RTA00000404F.i.18.1	21912	2	1	0	0	0	0	0	0
1139	RTA00000422F.i.14.1	39300	2	0	0	0	0	0	0	0
1140	RTA00000418F.m.14.1	75711	1	0	0	1	0	0	0	0
1141	RTA00000406F.o.12.1	37459	2	0	0	0	0	0	0	0
1143	RTA00000411F.a.07.1	74547	1	0	0	0	0	0	0	0
1144	RTA00000411F.c.02.1	72852	1	0	0	0	0	0	0	0
1146	RTA00000130A.h.16.1	80761	1	0	0	0	0	0	0	0
1147	RTA00000410F.p.23.1	73948	1	0	0	0	0	0	0	0
1148	RTA00000418F.m.24.1	77114	1	0	0	0	0	0 -	0	0
1150	RTA00000408F.j.19.2	73752	1	0	0	0	0	0	0	0
1152	RTA00000118A.d.17.1	81921	1	0	0	0	0	0	0	0
1153	RTA00000407F.b.04.1	63221	1	0	0	0	0	0	0	0
1154	RTA00000411F.e.07.1	65008	1	0	0	0	0	0	0	0
1156	RTA00000132A.c.11.1	87278	1	0	0	0	0	0	0	0
1157	RTA00000420F.e.16.1	63639	1	0	0	0	0	0	0	0
1159	RTA00000404F.b.11.1	39079	2	0	0	0	0	0	0	0
1160	RTA00000418F.k.17.1	75390	1	0	0	0	0	0	0	0
1161	RTA00000129A.k.12.1	79322	1	0	0	0	0	0	0	0
1162	RTA00000340R.m.07.1	78415	1	0	0	0	0	0	0	0
1163	RTA00000405F.d.14.1	35209	2	0	0	0	0	0	1	0
1164	RTA00000406F.f.11.1	38601	2	0	0	0	0	0	0	0
1165	RTA00000120A.h.5.1	80344	1	0	0	0	0	0	0	0
1167	RTA00000411F.g.06.1	66065	1	0	0	0	0	0	0	0
1168	RTA00000408F.d.16.1	76318	1	0	0	0	0	0	0	0
1171	RTA00000404F.c.19.1	39026	2	0	0	0	0	0	0	1
1173	RTA00000410F.a.01.1	73354	1	0	0	0	0	0	0	0
				612						

 $|\mathcal{S}_{ij}^{(k)}(x_i)| \leq |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)|^2 + |\mathcal{S}_{ij}^{(k)}(x_i)$ 

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones				
1174	RTA00000408F.h.08.1	74575	1	0	0	0	0	0	0	0
1175	RTA00000422F.b.16.1	17045	4	0	0	0	0	0	0	0
1176	RTA00000419F.f.10.1	66193	1	0	0	0	0	0	0	0
1177	RTA00000418F.l.04.1	74140	1	0	0	0	0	0	0	0
1178	RTA00000410F.a.16.1	73548	1	0	0	0	0	0	0	0
1179	RTA00000138A.e.13.1	79608	1	0	0	0	0	0	0	0
1180	RTA00000130A.b.5.1	79579	1	0	0	0	0	0	0	0
1181	RTA00000408F.j.15.2	74759	1	0	0	0	0	0	0	0
1182	RTA00000410F.m.20.1	74285	1	0	0	0	0	0	0	0
1185	RTA00000419F.e.04.1	62963	1	0	0	0	0	0	0	0
1187	RTA00000418F.g.05.1	73075	1	0	0	0	0	0	0	0
1188	RTA00000419F.n.02.1	65963	1	0	0	0	0	0	0	0
1191	RTA00000119A.m.15.1	80989	1	0	0	0	0	0	0	0
1194	RTA00000413F.g.23.1	40700	1	1	0	0	0	0	0	0
1195	RTA00000403F.a.18.1	75726	1	0	0	0	0	0	0	0
1196	RTA00000404F.m.20.2	39144	2	0	0	0	0	0	0	0
1199	RTA00000419F.h.04.1	65034	1	0	0	0	0	0	0	0
1200	RTA00000408F.d.12.1	75782	1	0	0	0	0	0	0	0
1201	RTA00000133A.m.19.2	80167	1	0	0	0	0	0	0	0
1206	RTA00000126A.o.22.1	81752	1	0	0	0	0	0	0	0
1207	RTA00000419F.n.13.1	66026	1	0	0	0	0	0	0	0
1208	RTA00000130A.h.13.1	80790	1	0	0	0	0	0	- 0	0
1212	RTA00000411F.m.19.1	74924	1	0	0	0	0	0	0	0
1214	RTA00000419F.k.06.1	78493	1	0	0	0	0	0	0	0
1216	RTA00000412F.d.16.1	26829	1	0	0	0	0	0	0	0
1217	RTA00000119A.j.23.1	79835	1	0	0	0	0	0	0	0
1219	RTA00000195AF.c.12.1	37582	2	0	0	0	0	0	0	0
1223	RTA00000423F.c.19.1	40472	2	0	0	0	0	0	0	0
1224	RTA00000405F.g.24.1	39076	2	0	0	0	0	0	0	0
1226	RTA00000419F.c.11.1	65504	1	0	0	0	0	0	0	0
1227	RTA00000135A.f.14.2	79969	1	0	0	0	0	0	0	0
1228	RTA00000403F.a.05.1	18808	1	1	0	0	0	0	0	0
1229	RTA00000405F.e.17.1	38662	2	0	0	0	0	0	0	0
1230	RTA00000411F.d.05.1	75812	1	0	0	0	0	0	0	0
1232	RTA00000418F.d.03.1	76824	1	0	0	0	0	0	0	0
1233	RTA00000418F.h.08.1	76401	1	0	0	0	0	0	0	0
1234	RTA00000418F.m.10.1	79110	1	0	0	0	0	0	0	0
1235	RTA00000411F.i.15.1	31612	1	1	0	0	0	0	0	0
1236	RTA00000413F.i.23.1	63073	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones				
1237	RTA00000411F.e.24.1	64781	1	0	0	0	0	0	0	0
1238	RTA00000406F.g.22.1	38590	2	0	0	0	0	0	0	0
1239	RTA00000126A.n.13.2	79735	1	0	0	0	0	0	0	0
1240	RTA00000419F.a.02.1	77993	1	0	0	0	0	0	0	0
1241	RTA00000346F.I.13.1	7542	8	0	0	2	1	0	1	0
1245	RTA00000120A.d.15.1	80533	1	0	0	0	0	0	0	0
1246	RTA00000418F.f.21.1	75157	1	0	0	0	0	0	0	0
1248	RTA00000129A.d.1.2	80058	1	0	0	0	0	0	0	0
1251	RTA00000419F.m.20.1	76720	1	0	0	0	0	0	0	0
1253	RTA00000406F.e.15.1	39074	2	0	0	0	0	0	0	0
1255	RTA00000411F.c.10.1	73117	1	0	0	0	0	0	0	0
1259	RTA00000413F.d.05.1	64788	1	0	0	0	0	0	0	0
1260	RTA00000121A.o.3.1	81437	1	0	0	0	0	0	0	0
1262	RTA00000420F.e.02.1	40259	2	0	0	0	0	0	0	0
1268	RTA00000126A.k.7.2	79866	1	0	0	0	0	0	0	0
1270	RTA00000419F.l.03.1	79060	1	0	0	0	0	0	0	0
1272	RTA00000118A.a.2.1	38067	2	0	0	0	0	0	0	0
1273	RTA00000410F.m.18.1	76365	1	0	0	0	0	0	0	0
1275	RTA00000406F.c.20.1	38578	2	0	0	0	0	0	0	0
1276	RTA00000413F.b.14.1	66591	1	0	0	0	0	0	0	0
1277	RTA00000406F.c.18.1	14368	2	0	0	0	0	0	0	0
1278	RTA00000418F.j.09.1	76352	1	0	0	0	0	0 -	0	0
1279	RTA00000419F.f.23.1	65002	1	0	0	0	0	0	0	0
1281	RTA00000411F.a.05.1	76699	1	0	0	0	0	0	0	0
1282	RTA00000419F.m.21.1	77947	1	0	0	0	0	0	0	0
1283	RTA00000405F.n.16.1	21503	2	1	1	0	0	0	0	0
1284	RTA00000422F.o.19.2	13084	3	2	0	0	0	0	0	0
1285	RTA00000408F.n.02.2	76993	1	0	0	0	0	0	0	0
1290	RTA00000119A.g.7.1	83580	1	0	0	0	0	0	0	0
1291	RTA00000411F.i.02.1	66975	1	0	0	0	0	0	0	0
1292	RTA00000408F.l.09.1	75487	1	0	0	0	0	0	0	0
1293	RTA00000423F.g.04.1	23012	2	1	0	0	0	0	0	0
1295	RTA00000418F.i.18.1	78024	1	0	0	0	0	0	0	0
1296	RTA00000411F.h.15.1	65160	1	0	0	0	0	0	0	0
1297	RTA00000410F.i.19.1	78988	1	0	0	0	0	0	0	0
1298	RTA00000419F.k.24.1	75596	1	0	0	0	0	0	0	0
1301	RTA00000409F.i.09.1	75279	1	0	0	0	0	0	0	0
1302	RTA00000419F.h.02.1	63985	1	0	0	0	0	0	0	0
1303	RTA00000413F.b.12.1	64932	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones		lib 16 clones			lib 19 clones	lib 20 clones
1304	RTA00000121A.h.18.1	16376	4	0	0	0	0	0	0	0
1305	RTA00000411F.n.20.1	75816	1	0	0	0	0	0	0	0
1307	RTA00000411F.n.12.1	73308	1	0	0	0	0	0	0	0
1308	RTA00000408F.j.12.2	18226	1	0	0	0	0	0	0	0
1309	RTA00000409F.i.03.1	75968	1	0	0	0	0	0	0	0
1312	RTA00000409F.j.05.1	74128	1	0	0	0	0	0	0	0
1313	RTA00000419F.m.04.1	74367	1	0	0	0	0	0	0	0
1314	RTA00000418F.k.03.1	78901	1	0	0	0	0	0	0	0
1315	RTA00000419F.d.16.1	64357	1	0	0	0	0	0	0	0
1316	RTA00000420F.e.10.1	65899	1	0	0	0	0	0	0	0
1319	RTA00000418F.k.08.1	18259	1	0	0	0	0	0	0	0
1322	RTA00000410F.c.02.1	75055	1	0	0	0	0	0	0	0
1324	RTA00000403F.h.18.1	39241	2	0	0	0	0	0	0	0
1325	RTA00000405F.n.13.1	23810	2	1	0	0	0	0	0	0
1326	RTA00000355R.e.14.1	16837	2	2	0	0	0	0	0	0
1327	RTA00000422F.1.03.1	39147	2	0	0	0	0	0	0	0
1329	RTA00000403F.o.14.1	38971	2	0	0	0	0	0	0	0
1333	RTA00000127A.f.11.1	81463	1	0	0	0	0	0	0	0
1335	RTA00000403F.o.07.1	39037	2	0	0	0	0	0	0	0
1336	RTA00000403F.d.19.1	39243	2	0	0	0	0	0	0	0
1338	RTA00000406F.i.17.1	37902	2	0	0	0	0	0	0	0
1339	RTA00000418F.d.22.1	75324	1	0	0	0	0	0 -	0	0
1340	RTA00000340R.o.12.1	53732	1	0	0	0	0	0	0	0
1341	RTA00000125A.g.24.1	80397	1	0	0	0	0	0	0	0
1342	RTA00000130A.o.21.1	80218	1	0	0	0	0	0	0	0
1343	RTA00000420F.a.23.1	42158	1	1	0	0	0	0	0	0
1344	RTA00000411F.m.18.1	75629	1	0	0	0	0	0	0	0
1345	RTA00000407F.b.22.1	37487	2	0	0	0	0	0	0	0
1346	RTA00000409F.a.16.1	73990	1	0	0	0	0	0	0	0
1348	RTA00000341F.k.12.1	62985	1	0	0	0	0	0	0	0
1349	RTA00000129A.c.18.2	37216	2	0	0	0	0	0	0	0
1350	RTA00000410F.d.10.1	77561	1	0	0	0	0	0	0	0
1351	RTA00000351R.i.03.1	6874	6	3	0	0	1	0	0	0
1352	RTA00000135A.I.1.2	39426	2	0	0	0	0	0	0	0
1353	RTA00000420F.b.18.1	66136	1	0	0	0	0	0	0	0
1356	RTA00000403F.o.13.1	39049	2	0	0	0	0	0	0	0
1357	RTA00000411F.f.06.1	64186	1	0	0	0	0	0	0	0
1359	RTA00000351R.c.13.1	11476	6	0	0	0	0	0	0	0
1362	RTA00000420F.d.16.1	64485	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones			lib 19 clones	
1363	RTA00000404F.i.12.1	39001	2	0	0	0	0	0	0	0
1364	RTA00000404F.o.10.2	16785	2	2	0	0	0	0	0	0
1365	RTA00000419F.d.07.1	21421	1	2	0	0	0	0	0	0
1366	RTA00000404F.p.02.2	39097	2	0	1	0	0	0	0	0
1367	RTA00000125A.k.14.1	79457	1	0	0	0	0	0	0	0
1368	RTA00000122A.j.22.1	81151	1	0	0	0	0	0	0	0
1369	RTA00000406F.i.13.1	37904	2	0	0	0	0	0	0	0
1370	RTA00000135A.b.23.1	35241	2	0	0	0	0	0	0	0
1373	RTA00000423F.1.04.1	14320	2	0	0	0	0	0	0	0
1374	RTA00000420F.b.04.1	63820	1	0	0	0	0	0	0	0
1376	RTA00000408F.i.18.2	74410	1	0	0	0	0	0	0	0
1378	RTA00000341F.j.05.1	36177	2	0	0	0	0	0	0	0
1379	RTA00000420F.a.16.1	63345	1	0	0	0	0	0	0	0
1381	RTA00000410F.j.01.1	73399	1	0	0	0	0	0	0	0
1382	RTA00000408F.p.21.1	77930	1	0	0	0	0	0	0	0
1383	RTA00000412F.d.19.1	75743	1	0	0	0	0	0	0	0
1384	RTA00000352R.c.04.1	71976	1	0	0	0	0	0	0	0
1385	RTA00000413F.f.19.1	65189	1	0	0	0	0	0	0	0
1386	RTA00000411F.e.03.1	73648	1	0	0	0	0	0	0	0
1389	RTA00000418F.c.04.1	41587	1	1	0	0	0	0	0	0
1390	RTA00000418F.o.17.1	79069	1	0	0	0	0	0	0	0
1391	RTA00000418F.e.21.1	74773	1	0	0	0	0	0 -	0	0
1392	RTA00000419F.d.14.1	64945	1	0	0	0	0	0	0	0
1396	RTA00000410F.j.20.1	73601	1	0	0	0	0	0	0	0
1399	RTA00000119A.j.9.1	82060	1	0	0	0	0	0	0	0
1403	RTA00000340F.i.13.1	79299	1	0	0	0	0	0	0	0
1404	RTA00000412F.g.03.1	64740	1	0	0	0	0	0	0	0
1405	RTA00000122A.g.17.1	32655	1	1	0	0	0	0	0	0
1407	RTA00000419F.n.12.1	66086	1	0	0	0	0	0	0	0
1410	RTA00000351R.p.14.1	13166	2	3	0	0	0	0	0	0
1411	RTA00000403F.e.08.1	19126	3	0	0	0	0	0	0	0
1412	RTA00000124A.k.20.1	80913	1	0	0	0	0	0	0	0
1413	RTA00000121A.n.2.1	33585	1	1	0	0	0	0	0	0
1414	RTA00000422F.m.24.1	39159	2	0	1	0	1	1	2	2
1415	RTA00000408F.e.24.2	75002	1	0	0	0	0	0	0	0
1418	RTA00000403F.b.12.1	78775	1	0	0	0	0	0	0	0
1419	RTA00000404F.a.09.1	38985	2	0	0	0	0	0	0	0
1421	RTA00000403F.o.19.1	78615	1	0	0	0	0	0	0	0
1424	RTA00000410F.b.10.1	74504	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones				lib 18 clones		
1426	RTA00000413F.h.12.1	66929	1	0	0	0	0	0	0	0
1427	RTA00000406F.k.14.1	38651	2	0	0	0	0	0	0	0
1429	RTA00000411F.f.17.1	65661	1	0	0	0	0	0	0	0
1430	RTA00000411F.k.10.1	64506	1	0	0	0	0	0	0	0
1431	RTA00000411F.g.21.1	64500	1	0	0	0	0	0	0	0
1432	RTA00000119A.h.24.1	82266	1	0	0	0	0	0	0	0
1434	RTA00000408F.m.22.2	72949	1	0	0	0	0	0	0	0
1437	RTA00000410F.i.17.1	78147	1	0	0	0	0	0	0	0
1440	RTA00000129A.a.13.2	79780	1	0	0	0	0	0	0	0
1441	RTA00000129A.k.21.1	82067	1	0	0	0	0	0	0	0
1442	RTA00000350R.g.10.1	9026	7	0	0	1	0	0	0	0
1443	RTA00000413F.d.23.1	66030	1	0	0	0	0	0	0	0
1447	RTA00000411F.d.10.1	76445	1	0	0	0	0	0	0	0
1448	RTA00000404F.b.19.1	39281	2	0	0	0	0	0	0	0
1449	RTA00000418F.c.07.1	73245	1	0	0	0	0	0	0	0
1450	RTA00000418F.j.15.1	74855	1	0	0	0	0	1	0	0
1453	RTA00000413F.b.16.1	65126	1	0	0	0	0	0	0	0
1455	RTA00000350R.m.14.1	39171	2	0	0	0	0	0	0	0
1456	RTA00000418F.I.11.1	77158	1	0	0	0	0	0	0	0
1457	RTA00000130A.d.5.1	82051	1	0	0	0	0	0	0	0
1458	RTA00000339F.n.05.1	39648	2	0	0	0	0	0	0	0
1460	RTA00000407F.a.23.1	23489	2	1	0	0	0	0 -	0	0
1462	RTA00000403F.h.11.1	39219	2	0	0	0	0	0	0	0
1463	RTA00000406F.j.13.1	38688	2	0	0	0	0	0	0	0
1464	RTA00000352R.p.09.1	16915	4	0	0	0	0	0	0	0
1465	RTA00000413F.g.24.1	65481	1	0	0	0	0	0	0	0
1469	RTA00000420F.a.08.1	19473	1	2	0	0	0	0	0	0
1472	RTA00000404F.i.22.1	39082	2	0	0	0	0	0	0	0
1473	RTA00000124A.k.23.1	81350	1	0	0	0	0	0	0	0
1474	RTA00000404F.e.11.1	38991	2	0	0	0	0	0	0	0
1475	RTA00000129A.d.2.4	80119	1	0	0	0	0	0	0	0
1478	RTA00000419F.o.15.1	32487	1	1	0	0	0	0	0	0
1479	RTA00000119A.m.17.1	79536	1	0	0	0	0	0	0	0
1480	RTA00000410F.b.07.1	78916	1	0	0	0	0	0	0	0
1481	RTA00000420F.b.19.1	36873	2	0	0	0	0	0	0	0
1483	RTA00000411F.b.21.1	10051	1	0	0	0	0	0	0	0
1485	RTA00000356R.c.16.1	16915	4	0	0	0	0	0	0	0
1487	RTA00000412F.h.11.1	63175	1	0	0	0	0	0	0	0
1490	RTA00000420F.a.11.1	66460	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones				
1491	RTA00000120A.c.7.1	80985	1	0	0	1	0	0	0	0
1492	RTA00000404F.e.15.1	39101	2	0	0	0	0	0	0	0
1493	RTA00000422F.n.20.1	38676	2	0	0	0	0	0	1	0
1494	RTA00000423F.h.20.1	38639	2	0	0	0	0	0	0	0
1497	RTA00000410F.b.18.1	76701	1	0	0	0	0	0	0	0
1499	RTA00000423F.g.15.1	35173	2	0	0	0	0	0	0	0
1500	RTA00000413F.b.04.1	66427	1	0	0	0	0	0	0	0
1503	RTA00000346F.f.11.1	38528	2	0	0	0	0	0	0	0
1506	RTA00000422F.i.02.1	76436	1	0	0	0	0	0	0	0
1507	RTA00000410F.a.08.1	73324	1	0	0	0	0	0	0	0
1509	RTA00000419F.e.02.1	65010	1	0	0	0	0	0	0	0
1511	RTA00000403F.g.13.1	38718	2	0	0	0	0	0	0	0
1513	RTA00000407F.a.01.1	12501	3	1	0	0	0	0	0	0
1516	RTA00000411F.f.14.1	62984	1	0	0	0	0	0	0	0
1517	RTA00000411F.c.04.1	76858	1	0	0	0	0	0	0	0
1518	RTA00000135A.m.18.1	19255	2	0	0	0	0	0	0	0
1519	RTA00000413F.c.17.1	36831	2	0	0	0	0	0	0	0
1521	RTA00000404F.j.01.1	26859	2	0	0	0	0	0	0	0
1522	RTA00000138A.p.10.1	81625	1	0	0	0	0	0	0	0
1526	RTA00000423F.h.07.1	37933	2	0	0	0	0	0	0	0
1527	RTA00000413F.e.04.1	64176	1	0	0	0	0	0	0	0
1528	RTA00000406F.h.03.1	38585	2	0	0	0	0	0 -	0	0
1529	RTA00000403F.e.24.1	16432	2	2	0	0	0	0	0	0
1531	RTA00000403F.i.11.1	23535	2	1	0	0	0	0	0	0
1532	RTA00000419F.g.02.1	62839	1	0	0	0	0	0	0	0
1533	RTA00000347F.e.05.1	39814	2	0	0	0	0	0	0	0
1534	RTA00000408F.l.16.1	73468	1	0	0	0	0	0	0	0
1536	RTA00000423F.f.09.1	64823	1	0	0	0	0	0	0	0
1537	RTA00000419F.k.03.1	40822	1	1	0	0	0	0	0	0
1538	RTA00000406F.b.02.1	38744	2	0	0	0	0	0	0	0
1539	RTA00000418F.o.14.1	33524	1	1	0	0	0	0	0	0
1541	RTA00000404F.b.09.1	39166	2	0	0	0	0	0	0	0
1547	RTA00000406F.k.11.1	38715	2	0	0	0	0	0	0	0
1549	RTA00000406F.c.06.1	37924	2	0	0	0	0	0	0	0
1550	RTA00000418F.n.07.1	76316	1	0	0	0	0	0	0	0
1551	RTA00000419F.n.15.1	63484	1	0	0	0	0	0	0	0
1552	RTA00000408F.n.06.2	76642	1	0	0	0	0	0	0	0
1553	RTA00000420F.c.04.1	65007	1	0	0	0	0	0	0	0
1554	RTA00000411F.j.15.1	66871	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones	lib 18 clones	lib 19 clones	lib 20 clones
1556	RTA00000128A.m.23.1	81441	1	0	0	0	0	0	0	0
1557	RTA00000406F.g.03.1	38690	2	0	0	0	0	0	0	0
1558	RTA00000405F.h.05.2	75706	1	0	0	0	0	0	0	0
1559	RTA00000129A.n.24.1	81409	1	0	0	0	0	0	0	0
1562	RTA00000418F.n.11.1	78977	1	0	0	0	0	0	0	0
1565	RTA00000120A.h.9.1	80736	1	0	0	0	0	0	0	0
1566	RTA00000413F.a.12.1	63403	1	0	0	0	0	0	0	0
1567	RTA00000412F.o.05.1	63575	1	0	0	0	0	0	0	0
1571	RTA00000354R.n.04.1	22049	3	0	0	0	0	0	0	0
1573	RTA00000406F.h.05.1	38542	2	0	0	0	0	0	0	0
1574	RTA00000410F.b.24.1	75104	1	0	0	0	0	0	0	0
1575	RTA00000423F.d.11.1	38950	2	0	0	0	0	0	0	0
1578	RTA00000119A.k.1.1	81282	1	0	0	0	0	0	0	0
1579	RTA00000420F.f.07.1	66312	1	0	0	0	0	0	0	0
1580	RTA00000404F.k.22.2	39084	2	0	0	0	0	0	0	0
1581	RTA00000422F.e.07.1	38964	2	0	0	0	0	0	0	0
1582	RTA00000410F.f.12.1	73883	1	0	0	0	0	0	0	0
1584	RTA00000411F.m.11.1	73196	1	0	0	0	0	0	0	0
1587	RTA00000403F.o.10.2	38964	2	0	0	0	0	0	0	0
1590	RTA00000413F.c.10.1	65600	1	0	0	0	0	0	0	0
1591	RTA00000411F.b.17.1	72893	1	0	0	0	0	0	0	0
1593	RTA00000408F.k.19.1	77593	1	0	0	0	0	0	0	0
1596	RTA00000119A.i.8.1	82593	1	0	0	0	0	0	0	0
1598	RTA00000418F.g.03.1	78737	1	0	0	0	0	0	0	0
1599	RTA00000411F.a.09.1	78629	1	0	0	0	0	0	0	0
1601	RTA00000419F.j.11.1	73183	1	0	0	0	0	0	0	0
1603	RTA00000404F.n.18.2	37169	2	0	0	0	0	0	0	0
1604	RTA00000122A.n.16.1	80553	1	0	0	0	0	0	0	0
1605	RTA00000420F.c.07.1	65555	1	0	0	0	0	0	0	0
1608	RTA00000408F.j.13.2	42275	1	1	0	0	0	0	0	0
1610	RTA00000423F.a.01.1	39103	2	0	0	0	0	0	0	0
1613	RTA00000341F.e.20.1	67422	1	0	0	0	0	0	0	0
1614	RTA00000419F.m.22.1	75600	1	0	0	0	0	0	0	0
1615	RTA00000419F.m.23.1	64263	1	0	0	0	0	0	0	0
1616	RTA00000419F.b.06.1	76728	1	0	0	0	0	0	0	0
1618	RTA00000406F.p.08.1	37573	2	<b>, 0</b>	0	0	0	0	0	2
1619	RTA00000129A.n.17.1	79811	1	0	0	0	0	0	0	0
1621	RTA00000407F.b.08.1	37513	2	0	0	0	0	0	0	0
1623	RTA00000406F.i.08.1	37946	2	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones	lib 18 clones	lib 19 clones	lib 20 clones
1624	RTA00000403F.h.07.1	26856	2	0	0	0	0	0	0	0
1625	RTA00000418F.n.24.1	73153	1	0	0	0	0	0	0	0
1627	RTA00000409F.l.20.1	74394	1	0	0	0	0	0	0	0
1628	RTA00000418F.l.06.1	73317	1	0	0	0	0	0	0	0
1629	RTA00000346F.o.22.1	7381	2	6	0	0	0	0	0	0
1630	RTA00000129A.k.22.1	79639	1	0	0	0	0	0	0	0
1632	RTA00000418F.m.22.1	74567	1	0	0	0	0	0	0	0
1633	RTA00000413F.c.12.1	65334	1	0	0	0	0	0	0	0
1635	RTA00000418F.g.20.1	74626	1	0	0	0	0	0	0	0
1636	RTA00000413F.d.15.1	64943	1	0	0	0	0	0	0	0
1639	RTA00000412F.c.10.1	76372	1	0	0	0	0	0	0	0
1640	RTA00000122A.j.17.1	62736	1	0	0	0	0	0	0	0
1645	RTA00000418F.j.19.1	78399	1	0	0	0	0	0	0	0
1646	RTA00000137A.p.12.1	80614	1	0	0	0	0	0	0	0
1648	RTA00000418F.p.10.1	75323	1	0	0	0	0	0	0	0
1649	RTA00000408F.k.12.1	77246	1	0	0	0	0	0	0	0
1650	RTA00000137A.j.11.4	79752	1	0	0	0	0	0	0	0
1652	RTA00000419F.n.24.1	65995	1	0	0	0	0	0	0	0
1653	RTA00000418F.1.03.1	79058	1	0	0	0	0	0	0	0
1655	RTA00000419F.m.13.1	79052	1	0	0	0	0	0	0	0
1656	RTA00000418F.j.14.1	32623	1	1	0	0	0	0	0	0
1657	RTA00000403F.a.10.1	73952	1	0	0	0	0	0	-0	0
1658	RTA00000420F.a.21.1	66241	1	0	0	0	0	0	0	0
1659	RTA00000127A.e.6.1	5885	4	2	0	0	0	0	0	0
1660	RTA00000405F.g.21.2	38966	2	0	0	0	0	0	0	0
1661	RTA00000405F.g.21.1	38966	2	0	0	0	0	0	0	0
1662	RTA00000419F.m.06.1	75749	1	0	0	0	0	0	0	0
1663	RTA00000423F.g.03.1	38007	2	0	0	0	0	0	0	0
1665	RTA00000418F.f.03.1	78911	1	0	0	0	0	0	0	0
1668	RTA00000120A.c.20.1	43235	1	1	0	0	0	1	0	0
1669	RTA00000138A.m.15.1	41603	1	1	0	0	0	0	0	0
1670	RTA00000408F.f.14.2	73024	1	0	0	0	0	0	0	0
1671	RTA00000418F.p.20.1	78023	1	0	0	0	0	0	0	0
1672	RTA00000423F.e.21.1	66961	1	0	0	0	0	0	0	0
1673	RTA00000419F.j.22.1	73525	1	0	0	0	0	0	0	0
1674	RTA00000410F.d.18.1	75458	1	0	0	0	0	0	0	0
1675	RTA00000403F.b.24.1	78838	1	0	0	0	0	0	0	0
1677		76093	1	0	0	0	0	0	0	0
1680	RTA00000353R.h.10.1	39498	2	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones	lib 18 clones	lib 19 clones	lib 20 clones
1682	RTA00000411F.d.21.1	74794	1	0	0	0	0	0	0	0
1683	RTA00000340F.m.04.1	19406	2	1	0	0	0	0	0	0
1684	RTA00000411F.n.09.1	78962	1	0	0	0	0	0	0	0
1685	RTA00000127A.h.22.2	13155	2	3	0	0	0	0	0	0
1686	RTA00000420F.e.09.1	66325	1	0	0	0	0	0	0	0
1687	RTA00000405F.p.03.1	11346	3	3	0	0	0	0	0	0
1688	RTA00000419F.a.18.1	78484	1	0	0	0	0	0	0	0
1691	RTA00000121A.n.23.1	26981	2	0	0	0	0	0	0	0
1692	RTA00000121A.n.15.1	40849	1	1	0	0	0	0	0	0
1693	RTA00000403F.i.23.1	11364	4	2	0	0	0	0	0	0
1694	RTA00000405F.a.03.1	39065	2	0	0	0	0	0	0	0
1696	RTA00000419F.p.08.1	65560	1	0	0	0	0	0	0	0
1697	RTA00000126A.n.6.2	79917	1	0	0	0	0	0	0	0
1698	RTA00000413F.c.03.1	64527	1	0	0	1	0	0	0	0
1699	RTA00000422F.k.24.1	39118	2	0	0	0	0	0	0	0
1700	RTA00000412F.c.17.1	75620	1	0	0	0	0	0	0	0
1702	RTA00000347F.g.08.1	23121	3	0	0	0	0	0	0	0
1703	RTA00000419F.o.06.1	64643	1	0	0	0	0	0	0	0
1704	RTA00000340R.j.07.1	38954	2	0	0	0	0	0	0	0
1705	RTA00000423F.j.02.1	38617	2	0	0	0	0	0	0	0
1706	RTA00000419F.c.04.1	63749	1	0	0	0	0	0	0	0
1707	RTA00000411F.a.01.1	74524	1	0	0	0	0	0	0	0
1708	RTA00000406F.f.05.1	22961	2	1	0	0	0	0	1	0
1709	RTA00000410F.n.05.1	77830	1	0	0	0	0	0	0	0
1710	RTA00000404F.e.06.1	39315	2	0	0	0	0	0	0	0
1712	RTA00000411F.c.03.1	79280	1	0	0	0	0	0	0	0
1718	RTA00000405F.l.07.1	38636	2	0	0	0	0	0	0	0
1720	RTA00000411F.n.06.1	73886	1	0	0	0	0	0	0	0
1721	RTA00000422F.k.15.1	19253	2	0	0	0	0	0	0	0
1722	RTA00000406F.h.16.1	38618	2	0	0	0	0	0	0	0
1723	RTA00000419F.f.24.1	18717	1	1	0	0	0	0	0	0
1724	RTA00000411F.d.18.1	76063	1	0	0	0	0	0	0	0
1727	RTA00000408F.d.15.1	78467	1	0	0	0	0	0	0	0
1728	RTA00000339F.b.22.1	6867	7	3	0	0	0	0	0	0
1730	RTA00000411F.n.02.1	78049	1	0	0	0	0	0	0	0
1731	RTA00000419F.b.17.1	63261	1	0	0	0	0	0	0	0
1733	RTA00000130A.e.20.1	79502	1	0	0	0	0	0	0	0
1735	RTA00000411F.i.13.1	66138	1	0	0	0	0	0	0	0
1736	RTA00000420F.e.20.1	64762	1	0	0	0	0	0	0	0

1737 RTA00000126A.p.23.2 80915 1 0 0 0 0 0 0 0 0 0 0 1 1739 RTA00000406F.g.08.1 37963 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1740 RTA00000406F.g.08.1 37963 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1741 RTA00000406F.d.24.1 37997 2 0 0 0 0 0 0 0 0 0 0 0 1 1744 RTA00000418F.i.12.1 78971 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1745 RTA00000418F.i.12.1 78971 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1746 RTA00000419F.b.10.1 78566 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1747 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1747 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1749 RTA00000406F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1749 RTA00000418F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000418F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000418F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1753 RTA00000418F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1755 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1755 RTA00000419F.g.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1755 RTA00000419F.g.24.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000418F.k.04.1 30053 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000418F.k.04.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000418F.k.03.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1770 RTA00000419F.g.23.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1777 RTA00000419F.g.03.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.g.03.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.g.03.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.g.03.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.g.03.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.g.03.1 38980 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.g.18.1 64071 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.g.18.1 64071 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.g.18.1 64071 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA000000419F.g.13.1 38980 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000040F.g.08.1 38980 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones		lib 16 clones				
1740 RTA00000409F.a.08.1 74978 1 0 0 0 0 0 0 0 0 0 1 1741 RTA00000406F.d.24.1 37997 2 0 0 0 0 0 0 0 0 0 0 0 1 1744 RTA00000418F.i.12.1 78971 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1745 RTA00000418F.i.12.1 78971 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1745 RTA00000419F.b.10.1 78566 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1746 RTA00000419F.b.10.1 78566 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1747 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1748 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1749 RTA00000406F.b.05.1 37894 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1749 RTA00000406F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1750 RTA00000419F.b.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000419F.b.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1753 RTA00000419F.b.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1755 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000419F.a.24.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000419F.a.24.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000419F.a.24.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000419F.a.24.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000419F.a.24.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		RTA00000126A.p.23.2	80915	1	0	0	0	0	0	0	0
1741 RTA00000406F.d.24.1 37997 2 0 0 0 0 0 0 0 0 0 1 1744 RTA00000418F.i.12.1 78971 1 0 0 0 0 0 0 0 0 0 0 0 1 1745 RTA00000121A.h.19.1 80334 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1746 RTA00000419F.b.10.1 78566 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1747 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1747 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1748 RTA00000406F.o.05.1 37894 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1749 RTA00000406F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1749 RTA00000406F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA000001314A.l.9.1 81814 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA0000018F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1754 RTA00000418F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000418F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000418F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1758 RTA00000408F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000408F.a.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000419F.b.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000419F.b.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1770 RTA00000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1771 RTA00000149F.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1772 RTA00000149F.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1774 RTA00000149F.b.6.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.b.6.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1778 RTA00000419F.b.6.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.8.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.8.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.8.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1739	RTA00000406F.g.08.1	37963	2	0	0	0	0	0	0	0
1744 RTA00000418F.i.12.1 78971 1 0 0 0 0 0 0 0 0 0 0 1 1745 RTA00000419F.b.10.1 78566 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1746 RTA00000419F.b.10.1 78566 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1747 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1748 RTA00000406F.b.05.1 37894 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1748 RTA00000406F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1750 RTA0000041F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA0000041F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA0000041F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1754 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1761 RTA0000019A.e.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000419F.a.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA0000040F.a.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000408F.n.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000408F.n.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1770 RTA00000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1771 RTA000001408F.a.03.1 73442 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1774 RTA00000419F.p.06.1 65485 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.b.01.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 1 1778 RTA00000419F.b.01.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 1 1778 RTA00000419F.b.01.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.01.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.c.13.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.c.13.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1740	RTA00000409F.a.08.1	74978	1	0	0	0	0	0	0	0
1745 RTA00000121A.h.19.1 80334 1 0 0 0 0 0 0 0 0 0 0 0 1 1746 RTA00000419F.b.10.1 78566 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1747 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1748 RTA00000406F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1748 RTA00000408F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1750 RTA00000411F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000134A.l.9.1 81814 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1754 RTA00000419F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000419F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1759 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000409F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000409F.a.16.1 19251 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000409F.a.16.2 373720 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000409F.a.12.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1770 RTA00000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1771 RTA00000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1771 RTA00000149F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1778 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1741	RTA00000406F.d.24.1	37997	2	0	0	0	0	0	0	0
1746 RTA0000419F.b.10.1 78566 1 0 0 0 0 0 0 0 0 0 0 1 1747 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 1 1748 RTA00000406F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 1 1749 RTA00000408F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 1 1750 RTA00000411F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000141F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1755 RTA00000149F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000149F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 1 1759 RTA00000149F.p.18.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000419F.a.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000404F.a.01.1 19251 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000408F.n.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1769 RTA00000412F.l.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1770 RTA00000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 1 1771 RTA00000406F.n.12.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1772 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 1 1774 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1779 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1779 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000423F.i.6.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000423F.i.6.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1788 RTA00000423F.i.6.1 38604 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1786 RTA00000404F.b.01.1 18225 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1790 RTA00000404F.b.01.1 18225 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1744	RTA00000418F.i.12.1	78971	1	0	0	0	0	0	0	0
1747 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 1 1748 RTA00000406F.m.10.1 38004 2 0 0 0 0 0 0 0 0 0 0 0 1 1748 RTA00000406F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1750 RTA00000411F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000134A.1.9.1 81814 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000418F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1759 RTA00000419F.a.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000419F.a.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000409F.a.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000409F.n.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1769 RTA00000409F.a.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 1 1770 RTA0000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 1 1771 RTA00000409F.a.12.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 1 1772 RTA00000418F.e.03.1 73442 1 0 0 0 0 0 0 0 0 0 0 0 1 1774 RTA00000403F.g.03.1 23537 2 1 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 1 1776 RTA0000043F.p.06.1 65485 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1778 RTA0000043F.g.03.1 23537 2 1 0 0 0 0 0 0 0 0 0 0 0 1 1778 RTA0000043F.g.16.1 64773 1 0 0 0 0 0 0 0 0 0 0 0 1 1781 RTA0000043F.g.16.1 64773 1 0 0 0 0 0 0 0 0 0 0 0 1 1782 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 1 1784 RTA0000043F.g.03.1 38604 2 0 0 0 0 0 0 0 0 0 0 0 1 1785 RTA0000041F.f.04.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1786 RTA00000449F.g.08.1 38980 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1787 RTA00000449F.g.08.1 38980 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1787 RTA00000449F.b.21.1 63366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1745	RTA00000121A.h.19.1	80334	1	0	0	0	0	0	0	0
1748 RTA00000406F.o.05.1 37894 2 0 0 0 0 0 0 0 0 0 0 1 1749 RTA00000408F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1750 RTA00000411F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000134A.1.9.1 81814 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1754 RTA00000418F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1754 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1755 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1755 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000419F.a.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA0000040F.a.01.1 19251 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000408F.n.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000408F.n.16.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1770 RTA0000408F.n.16.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1771 RTA0000408F.n.16.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1772 RTA00000418F.e.03.1 73442 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1774 RTA0000403F.g.03.1 23537 2 1 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000418F.e.03.1 73442 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1776 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1778 RTA00000419F.h.8.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1781 RTA0000043F.g.03.1 38604 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1784 RTA0000043F.g.03.1 38604 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1785 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1786 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1786 RTA0000041F.f.04.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1786 RTA00000449F.g.08.1 38980 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1787 RTA00000449F.b.15.1 18225 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1746	RTA00000419F.b.10.1	78566	1	0	0	0	0	0	0	0
1749 RTA00000408F.b.04.2 39933 2 0 0 0 0 0 0 0 0 0 0 0 1750 RTA00000411F.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 0 1752 RTA00000134A.l.9.1 81814 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1754 RTA00000418F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1757 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1757 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1761 RTA00000129A.e.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1747	RTA00000406F.m.10.1	38004	2	0	0	0	0	0	0	0
1750 RTA0000041IF.k.04.1 65407 1 0 0 0 0 0 0 0 0 0 0 0 1 1752 RTA00000134A.I.9.1 81814 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1754 RTA00000418F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1759 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1759 RTA00000419F.a.24.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1761 RTA00000129A.e.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000404F.a.01.1 19251 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000404F.a.11.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1769 RTA00000412F.l.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1770 RTA00000412F.l.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1771 RTA00000412F.a.03.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1772 RTA00000418F.e.03.1 73442 1 0 0 0 0 0 0 0 0 0 0 0 1 1772 RTA00000412F.p.06.1 65485 1 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000412F.p.06.1 65485 1 0 0 0 0 0 0 0 0 0 0 0 1 1779 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 1 1781 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 1 1781 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1785 RTA0000041F.f.04.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1785 RTA0000041F.f.04.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1748	RTA00000406F.o.05.1	37894	2	0	0	0	0	0	0	0
1752 RTA00000134A.I.9.1	1749	RTA00000408F.b.04.2	39933	2	0	0	0	0	0	0	0
1754 RTA00000418F.k.04.1 75864 1 0 0 0 0 0 0 0 0 0 0 0 1 1757 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 0 1 1759 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 1 1761 RTA00000129A.e.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1762 RTA00000404F.a.01.1 19251 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000404F.a.01.1 19251 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1765 RTA00000408F.n.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1769 RTA00000412F.l.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1770 RTA00000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1771 RTA00000406F.n.12.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1772 RTA00000418F.e.03.1 73442 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1772 RTA00000418F.e.03.1 73442 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000418F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1776 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1781 RTA00000419F.h.81.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1781 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1750	RTA00000411F.k.04.1	65407	1	0	0	0	0	0	0	0
1757 RTA00000419F.p.18.1 63002 1 0 0 0 0 0 0 0 0 0 0 1 759 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1752	RTA00000134A.l.9.1	81814	1	0	0	0	0	0	0	0
1759 RTA00000419F.a.24.1 79290 1 0 0 0 0 0 0 0 0 0 0 1761 RTA00000129A.e.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1754	RTA00000418F.k.04.1	75864	1	0	0	0	0	0	0	0
1761 RTA00000129A.e.14.1 80053 1 0 0 0 0 0 0 0 0 0 0 1762 RTA00000404F.a.01.1 19251 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1765 RTA00000408F.n.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 1769 RTA00000412F.1.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1770 RTA00000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1771 RTA00000406F.n.12.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 1772 RTA00000406F.n.12.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1757	RTA00000419F.p.18.1	63002	1	0	0	0	0	0	0	0
1762 RTA00000404F.a.01.1 19251 2 0 0 0 0 0 0 0 0 0 0 1765 RTA00000408F.n.16.2 73720 1 0 0 0 0 0 0 0 0 0 0 0 0 1769 RTA00000412F.l.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1759	RTA00000419F.a.24.1	79290	1	0	0	0	0	0	0	0
1765 RTA00000408F.n.16.2 73720 1 0 0 0 0 0 0 0 0 0 1769 RTA00000412F.l.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1761	RTA00000129A.e.14.1	80053	1	0	0	0	0	0	0	0
1769 RTA00000412F.1.14.1 62792 1 0 0 0 0 0 0 0 0 0 0 1 770 RTA00000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1762	RTA00000404F.a.01.1	19251	2	0	0	0	0	0	0	0
1770 RTA00000129A.b.6.2 39111 2 0 0 0 0 0 0 0 0 0 0 1 771 RTA00000406F.n.12.1 37517 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1765	RTA00000408F.n.16.2	73720	1	0	0	0	0	0	0	0
1771 RTA00000406F.n.12.1 37517 2 0 0 0 0 0 0 0 0 0 0 1 1772 RTA00000418F.e.03.1 73442 1 0 0 0 0 0 0 0 0 0 0 0 0 1 1774 RTA00000403F.g.03.1 23537 2 1 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000412F.p.06.1 65485 1 0 0 0 0 0 0 0 0 0 0 0 1 1776 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 1 1779 RTA00000351R.j.16.1 64773 1 0 0 0 0 0 0 0 0 0 0 0 1 1781 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 1 1782 RTA00000423F.i.16.1 38604 2 0 0 0 0 0 0 0 0 0 0 0 1 1784 RTA00000419F.f.04.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 1 1785 RTA00000125A.c.17.1 80619 1 0 0 0 0 0 0 0 0 0 0 1 1786 RTA00000423F.c.13.1 39059 2 0 0 0 0 0 0 0 0 0 0 1 1787 RTA00000423F.c.13.1 39059 2 0 0 0 0 0 0 0 0 0 0 1 1790 RTA00000404F.k.15.1 18225 2 0 0 0 0 0 0 0 0 0 0 1 1792 RTA00000339F.l.12.1 7711 4 1 0 0 0 0 0 0 0 0 0 1 1793 RTA00000406F.b.01.1 39006 2 0 0 0 0 0 0 0 0 0 0 0 1 1794 RTA00000407F.c.08.1 37549 2 0 0 0 0 0 0 0 0 0 0 0 1 1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 0 0 1 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1769	RTA00000412F.1.14.1	62792	1	0	0	0	0	0	0	0
1772 RTA00000418F.e.03.1 73442 1 0 0 0 0 0 0 0 0 0 0 0 1 1774 RTA00000403F.g.03.1 23537 2 1 0 0 0 0 0 0 0 0 0 0 1 1775 RTA00000412F.p.06.1 65485 1 0 0 0 0 0 0 0 0 0 0 0 1 1776 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 1 1779 RTA00000351R.j.16.1 64773 1 0 0 0 0 0 0 0 0 0 0 0 1 1781 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 1 1782 RTA00000423F.i.16.1 38604 2 0 0 0 0 0 0 0 0 0 0 0 1784 RTA00000411F.f.04.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 1 1785 RTA00000425F.c.17.1 80619 1 0 0 0 0 0 0 0 0 0 0 0 0 1786 RTA00000404F.g.08.1 38980 2 0 0 0 0 0 0 0 0 0 0 1787 RTA00000404F.k.15.1 18225 2 0 0 0 0 0 0 0 0 0 0 0 1790 RTA00000404F.k.15.1 18225 2 0 0 0 0 0 0 0 0 0 0 0 1792 RTA00000339F.l.12.1 7711 4 1 0 0 0 0 0 0 0 0 0 0 1793 RTA00000406F.b.01.1 39006 2 0 0 0 0 0 0 0 0 0 0 1794 RTA00000407F.c.08.1 37549 2 0 0 0 0 0 0 0 0 0 0 0 1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 0 0 0 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1770	RTA00000129A.b.6.2	39111	2	0	0	0	0	0	0	0
1774 RTA00000403F.g.03.1 23537 2 1 0 0 0 0 0 0 0 0 1 775 RTA00000412F.p.06.1 65485 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1771	RTA00000406F.n.12.1	37517	2	0	0	0	0	0	0	0
1775 RTA00000412F.p.06.1 65485 1 0 0 0 0 0 0 0 0 0 0 1 1776 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 0 1 1779 RTA00000351R.j.16.1 64773 1 0 0 0 0 0 0 0 0 0 0 0 1 1781 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 1 1782 RTA00000423F.i.16.1 38604 2 0 0 0 0 0 0 0 0 0 0 0 1 1784 RTA00000411F.f.04.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 1 1785 RTA00000125A.c.17.1 80619 1 0 0 0 0 0 0 0 0 0 0 0 1 1786 RTA00000404F.g.08.1 38980 2 0 0 0 0 0 0 0 0 0 0 0 1 1787 RTA00000423F.c.13.1 39059 2 0 0 0 0 0 0 0 0 0 0 0 0 1 1790 RTA00000404F.k.15.1 18225 2 0 0 0 0 0 0 0 0 0 0 0 1 1792 RTA00000339F.l.12.1 7711 4 1 0 0 0 0 0 0 0 0 0 0 1 1793 RTA00000406F.b.01.1 39006 2 0 0 0 0 0 0 0 0 0 0 0 1 1794 RTA00000407F.c.08.1 37549 2 0 0 0 0 0 0 0 0 0 0 1 1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 0 0 1 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0 0 0 0 0 0	1772	RTA00000418F.e.03.1	73442	1	0	0	0	0	0	0	0
1776 RTA00000419F.b.21.1 65366 1 0 0 0 0 0 0 0 0 0 0 1 1 7 9 RTA00000351R.j.16.1 64773 1 0 0 0 0 0 0 0 0 0 0 0 0 0 1 1 8 1 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1774	RTA00000403F.g.03.1	23537	2	1	0	0	0	0	0	0
1779 RTA00000351R.j.16.1 64773 1 0 0 0 0 0 0 0 0 0 0 1781 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1775	RTA00000412F.p.06.1	65485	1	0	0	0	0	0	0	0
1781 RTA00000419F.f.18.1 64047 1 0 0 0 0 0 0 0 0 0 0 1782 RTA00000423F.i.16.1 38604 2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1776	RTA00000419F.b.21.1	65366	1	0	0	0	0	0	0	0
1782 RTA00000423F.i.16.1 38604 2 0 0 0 0 0 0 0 0 0 0 1784 RTA00000411F.f.04.1 64526 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1779	RTA00000351R.j.16.1	64773	1	0	0	0	0	0	0	0
1784 RTA00000411F.f.04.1 64526 1 0 0 0 0 0 0 0 0 0 0 1785 RTA00000125A.c.17.1 80619 1 0 0 0 0 0 0 0 0 0 0 0 0 1786 RTA00000404F.g.08.1 38980 2 0 0 0 0 0 0 0 0 0 0 0 0 1787 RTA00000423F.c.13.1 39059 2 0 0 0 0 0 0 0 0 0 0 0 0 1790 RTA00000404F.k.15.1 18225 2 0 0 0 0 0 0 0 0 0 0 0 1792 RTA00000339F.l.12.1 7711 4 1 0 0 0 0 0 0 0 0 0 1793 RTA00000406F.b.01.1 39006 2 0 0 0 0 0 0 0 0 0 0 1794 RTA00000407F.c.08.1 37549 2 0 0 0 0 0 0 0 0 0 0 1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 0 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0 0 0 0	1781	RTA00000419F.f.18.1	64047	1	0	0	0	0	0	0	0
1785       RTA00000125A.c.17.1       80619       1       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0	1782	RTA00000423F.i.16.1	38604	2	0	0	0	0	0	0	0
1786 RTA00000404F.g.08.1 38980 2 0 0 0 0 0 0 0 0 0 1787 RTA00000423F.c.13.1 39059 2 0 0 0 0 0 0 0 0 0 0 1790 RTA00000404F.k.15.1 18225 2 0 0 0 0 0 0 0 0 0 0 0 1792 RTA00000339F.l.12.1 7711 4 1 0 0 0 0 0 0 0 0 0 1793 RTA00000406F.b.01.1 39006 2 0 0 0 0 0 0 0 0 0 0 1794 RTA00000407F.c.08.1 37549 2 0 0 0 0 0 0 0 0 0 1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0 0 0	1784	RTA00000411F.f.04.1	64526	1	0	0	0	0	0	0	0
1787       RTA00000423F.c.13.1       39059       2       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0       0	1785	RTA00000125A.c.17.1	80619	1	0	0	0	0	0	0	0
1790 RTA00000404F.k.15.1 18225 2 0 0 0 0 0 0 0 0 0 0 1792 RTA00000339F.l.12.1 7711 4 1 0 0 0 0 0 0 0 0 0 1793 RTA00000406F.b.01.1 39006 2 0 0 0 0 0 0 0 0 0 0 1794 RTA00000407F.c.08.1 37549 2 0 0 0 0 0 0 0 0 0 0 1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 0 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0 0 0	1786	RTA00000404F.g.08.1	38980	2	0	0	0	0	0	0	0
1792 RTA00000339F.l.12.1 7711 4 1 0 0 0 0 0 0 0 0 1793 RTA00000406F.b.01.1 39006 2 0 0 0 0 0 0 0 0 0 1794 RTA00000407F.c.08.1 37549 2 0 0 0 0 0 0 0 0 0 1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 0 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0 0	1787	RTA00000423F.c.13.1	39059	2	0	0	0	0	0	0	0
1793 RTA00000406F.b.01.1 39006 2 0 0 0 0 0 0 0 0 0 1794 RTA00000407F.c.08.1 37549 2 0 0 0 0 0 0 0 0 0 1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0 0	1790	RTA00000404F.k.15.1	18225	2	0	0	0	0	0	0	0
1794 RTA00000407F.c.08.1 37549 2 0 0 0 0 0 0 0 0 0 1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0	1792	RTA00000339F.1.12.1	7711	4	1	0	0	0	0	0	0
1796 RTA00000403F.b.05.1 74300 1 0 0 0 0 0 0 0 0 1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0 0	1793	RTA00000406F.b.01.1	39006	2	0	0	0	0	0	0	0
1800 RTA00000408F.j.05.2 73878 1 0 0 0 0 0 0	1794	RTA00000407F.c.08.1	37549	2	0	0	0	0	0	0	0
•	1796	RTA00000403F.b.05.1	74300	1	0	0	0	0	0	0	0
1802 RTA00000419F.c.14.1 65727 1 0 0 0 0 0 0	1800	RTA00000408F.j.05.2	73878	1	0	0	0	0	0	0	0
	1802	RTA00000419F.c.14.1	65727	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones		lib 16 clones				
1806	RTA00000346F.h.24.1	4379	9	2	0	0	0	0	0	0
1807	RTA00000420F.b.02.1	64013	1	0	0	0	0	0	0	0
1808	RTA00000413F.b.24.1	65117	1	0	0	0	0	0	0	0
1809	RTA00000412F.d.08.1	75328	1	0	0	0	0	0	0	0
1811	RTA00000419F.m.18.1	76014	1	0	0	0	0	0	0	0
1812	RTA00000419F.1.24.1	74628	1	0	0	0	0	0	0	0
1813	RTA00000408F.c.06.1	78619	1	0	0	0	0	0	0	0
1814	RTA00000405F.h.21.2	39072	2	0	0	0	0.	0	0	0
1816	RTA00000405F.g.05.2	38987	2	0	0	0	0	0	0	0
1817	RTA00000411F.f.20.1	63501	1	0	0	0	0	0	0	0
1819	RTA00000420F.d.19.1	43146	1	1	0	0	0	0	0	0
1820	RTA00000195R.a.06.1	35265	2	0	1	0	0	0	0	0
1821	RTA00000123A.f.2.1	80379	1	0	0	0	0	0	0	0
1822	RTA00000411F.j.11.1	66154	1	0	0	0	0	0	0	0
1827	RTA00000419F.j.03.1	77578	1	0	0	0	0	0	0	0
1829	RTA00000423F.h.11.1	38977	2	0	0	0	0	0 .	0	0
1830	RTA00000413F.b.17.1	21704	1	2	0	0	0	0	0	0
1833	RTA00000423F.f.03.1	63852	1	0	0	0	0	0	0	0
1834	RTA00000419F.e.10.1	63225	1	0	0	0	0	0	0	0
1836	RTA00000403F.d.02.1	39224	2	0	0	0	0	0	0	0
1838	RTA00000418F.j.20.1	77101	1	0	0	0	0	0	0	0
1846	RTA00000356R.h.05.1	35052	2	0	1	0	0	0 -	0	0
1848	RTA00000340F.i.15.1	26815	1	0	0	0	0	0	0	0
1850	RTA00000345F.c.12.1	23824	2	1	0	0	0	0	0	0
1852	RTA00000412F.o.03.1	65039	1	0	0	0	0	0	0	0
1853	RTA00000409F.d.16.1	76090	1	0	0	0	0	0	0	0
1856	RTA00000408F.j.17.2	78935	1	0	0	0	0	0	0	0
1857	RTA00000126A.j.15.2	40425	2	0	0	0	0	0	0	0
1861	RTA00000410F.b.17.1	77458	1	0	0	0	0	0	0	0
1862	RTA00000419F.l.22.1	78444	1	0	0	0	0	0	0	0
1864	RTA00000422F.f.22.1	38703	2	0	0	0	0	0	0	0
1867	RTA00000418F.c.05.1	76475	1	0	0	0	0	0	0	0
1868	RTA00000418F.p.21.1	78068	1	0	0	0	0	0	0	0
1870	RTA00000340F.i.08.1	12005	2	1	0	0	0	0	0	0
1871	RTA00000410F.o.04.1	79018	1	0	0	0	0	0	0	0
1872	RTA00000411F.l.16.1	16122	1	3	0	0	0	0	0	0
1873	RTA00000411F.j.03.1	66263	1	0	0	0	0	0	0	0
1874	RTA00000126A.k.24.1	39428	2	0	0	0	0	0	0	0
1876	RTA00000120A.m.10.3	81376	1	0	0	0	0	0	0	0

SEQ ID	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones	lib 18 clones	lib 19 clones	lib 20 clones
NO: 1877	RTA00000419F.f.16.1	64679	1	0	0	0	0	0	0	0
1878	RTA00000419F.1.10.1	42261	1	1	0	0	0	0	0	0
1881	RTA00000136A.h.6.1	81620	1	0	0	0	0	0	0	0
1886	RTA000001307.ii.o.1	73741	1	0	0	0	0	0	0	0
1888	RTA00000415F.1.03.1	38580	2	0	0	0	0	0	0	0
1889	RTA00000418F.m.02.1	74550	1	0	0	0	0	0	0	0
1891	RTA00000406F.c.05.1	22077	3	0	1	0	0	0	0	0
1893	RTA00000411F.k.21.1	65349	1	0	0	0	0	0	0	0
1897	RTA00000418F.i.06.1	75151	1	0	0	0	0	0	0	0
1898	RTA00000423F.a.03.1	26796	2	0	0	0	0	0	0	0
1900	RTA00000423F.k.21.2	37499	2	0	0	0	0	0	0	0
1902	RTA00000404F.c.18.1	38982	2	0	0	0	0	0	0	0
1905	RTA00000411F.g.24.1	65233	1	0	0	0	0	0	0	0
1907	RTA00000405F.m.07.1	37733	2	0	0	0	0	0	0	0
1908	RTA00000411F.j.07.1	66963	1	0	0	0	0	0	0	0
1910	RTA00000353R.h.04.1	17123	4	0	0	0	0	0	0	0
1911	RTA00000408F.f.10.2	75309	1	0	0	0	0	0	0	0
1913	RTA00000405F.o.03.1	37575	2	0	0	0	0	0	0	0
1914	RTA00000413F.b.18.1	39873	2	0	0	0	0	0	0	0
1920	RTA00000408F.c.08.1	73473	1	0	0	0	0	0	0	0
1922	RTA00000410F.c.06.1	77784	1	0	0	0	1	0	0	0
1924	RTA00000405F.b.08.1	39182	2	0	0	0	0	0	- 0	0
1925	RTA00000409F.1.24.1	73174	1	0	0	0	0	0	0	0
1926	RTA00000406F.j.06.1	38952	2	0	0	0	0	0	0	0
1927	RTA00000423F.h.03.1	37903	2	0	0	0	0	0	0	0
1929	RTA00000121A.k.22.1	79523	1	0	0	0	0	0	0	0
1931	RTA00000411F.m.06.1	24195	2	1	0	0	0	0	0	0
1932	RTA00000126A.b.9.1	81279	1	0	0	0	0	0	0	0
1935	RTA00000404F.1.05.1	38671	2	0	0	0	0	0	0	0
1941	RTA00000419F.p.10.1	41448	1	1	0	0	0	0	0	0
1942		81016	1	0	0	0	0	0	0	0
1948		63987		0	0	0	0	0	0	0
1949		63908		0	0	0	0	0	0	0
1952		80085	1	0	0	0	0	0	0	0
1953		65405		0	0	0	0	0	0	0
1955		38955		0	0	0	0	0	0	0
1957		22779		0	0	0	0	0	0	0
1959		75736		0	0	0	0	0	0	0
1961	RTA00000418F.b.20.1	73560	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones		lib 18 clones	lib 19 clones	lib 20 clones
1964	RTA00000408F.n.05.2	77883	1	0	0	0	0	0	0	0
1965	RTA00000419F.o.09.1	66396	1	0	0	0	0	0	0	0
1970	RTA00000422F.o.08.2	26832	2	0	0	0	0	0	0	0
1973	RTA00000418F.m.18.1	76479	1	0	0	0	0	0	0	0
1974	RTA00000347F.e.20.1	39911	2	0	0	0	0	0	0	0
1975	RTA00000419F.e.23.1	65772	1	0	0	0	0	0	0	0
1982	RTA00000411F.g.05.1	64664	1	0	0	0	0	0	0	0
1983	RTA00000404F.h.10.1	37148	2	0	0	0	0	0	0	0
1984	RTA00000422F.n.14.1	26787	2	0	0	0	0	0	0	0
1986	RTA00000120A.m.13.3	80608	1	0	0	0	0	0	0	0
1987	RTA00000412F.i.03.1	65617	1	0	0	0	0	0	0	0
1988	RTA00000418F.l.02.1	39316	2	0	0	0	0	0	0	0
1990	RTA00000411F.j.04.1	66219	1	0	0	0	0	0	0	0
1995	RTA00000404F.a.18.1	36267	2	0	0	0	0	0	0	0
1996	RTA00000408F.1.14.1	12001	2	3	0	0	0	0	0	0
1997	RTA00000405F.d.10.1	39000	2	0	0	0	0	0	0	0
1999	RTA00000418F.h.23.1	75153	1	0	0	0	0	0	0	0
2001	RTA00000418F.j.11.1	73853	1	0	0	0	0	0	0	0
2002	RTA00000408F.o.13.1	74895	1	0	0	0	0	0	0	0
2003	RTA00000419F.o.07.1	14059	1	0	0	0	0	0	0	0
2004	RTA00000419F.n.17.1	63186	1	0	0	0	0	0	0	0
2005	RTA00000403F.f.15.1	22768	3	0	0	0	0	0 -	0	0
2006	RTA00000408F.d.03.1	22768	3	0	0	0	0	0	0	0
2008	RTA00000346F.f.02.1	62757	1	0	0	0	0	0	0	0
2010	RTA00000413F.i.21.1	64066	1	0	0	0	0	0	0	0
2012	RTA00000419F.h.21.1	64828	1	0	0	0	0	0	0	0
2021	RTA00000121A.a.2.1	81843	1	0	0	0	0	0	0	0
2022	RTA00000527F.g.13.1	36035	2	0	0	0	0	0	0	0
2025	RTA00000426F.h.11.1	75479	1	0	0	0	0	0	0	0
2030	RTA00000522F.b.22.1	75181	1	0	0	0	0	0	0	0
2033	RTA00000522F.a.23.1	38613	2	0	0	0	0	0	0	0
2035	RTA00000523F.b.02.1	65163	1	0	0	0	0	0	0	0
2036	RTA00000425F.j.14.1		1	0	0	0	0	0	0	0
2039	RTA00000522F.e.16.1	75283	1	0	0	0	0	0	0	0
2042	RTA00000523F.h.17.1	65586	1	0	0	0	0	0	0	0
2044	RTA00000522F.p.07.1	76888	1	0	0	0	0	0	0	0
	RTA00000522F.n.08.1		1	0	0	0	0	0	0	0
2046	RTA00000425F.c.06.1		1	0	0	0	0	0	0	0
2047	RTA00000427F.b.23.1	64297	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones			lib 19 clones	
2048	RTA00000527F.p.02.1	36844	2	0	0	0	0	0	0	0
2049	RTA00000427F.d.08.1	63967	1	0	0	0	0	0	0	0
2051	RTA00000426F.m.07.1	63504	1	0	0	0	0	0	0	0
2052	RTA00000427F.c.10.1	65478	1	0	0	0	0	0	0	0
2055	RTA00000424F.m.15.1	73759	1	0	0	0	0	0	0	0
2056	RTA00000426F.f.11.1	63102	1	0	0	0	0	0	0	0
2058	RTA00000426F.f.20.1	65134	1	0	0	0	0	0	0	0
2063	RTA00000527F.i.19.2	38089	2	0	0	0	0	0	0	0
2068	RTA00000523F.e.18.1	62898	1	0	0	0	0	0	0	0
2069	RTA00000527F.k.21.1	36051	2	0	0	0	0	0	0	0
2072	RTA00000522F.n.02.1	74959	1	0	0	0	0	0	0	0
2075	RTA00000425F.f.19.1	32635	1	1	0	0	0	0	0	0
2076	RTA00000528F.e.23.1	19242	3	0	0	0	0	0	0	0
2077	RTA00000522F.n.16.1	26769	1	0	0	0	0	0	0	0
2078	RTA00000427F.c.20.1	26527	1	0	0	0	0	0	0	0
2079	RTA00000527F.k.06.1	12469	3	1	0	0	0	0	0	0
2081	RTA00000523F.i.06.1	66341	1	0	0	0	0	0	0	0
2082	RTA00000427F.f.21.1	36853	2	0	0	0	0	. 0	0	0
2083	RTA00000427F.j.19.1	41395	1	1	0	0	0	0	0	0
2084	RTA00000522F.b.01.1	75691	1	0	0	0	0	0	0	0
2085	RTA00000424F.i.24.1	79101	1	0	0	0	0	0	0	0
2086	RTA00000523F.c.01.1	65710	1	0	0	0	0	0	.0	0
2087	RTA00000427F.b.15.1	66891	1	0	0	0	0	0	0	0
2090	RTA00000522F.j.15.2	76535	1	0	0	. 0	0	0	0	0
2093	RTA00000426F.f.19.1	66701	1	0	1	0	0	0	0	0
2096	RTA00000523F.i.22.1	64688	1	0	0	0	0	0	0	0
2098	RTA00000425F.i.17.1	43213	1	1	0	0	0	0	0	0
2101	RTA00000425F.p.12.1	73219	1	0	0	0	0	0	0	0
2102	RTA00000427F.j.07.1	64819	1	0	0	0	0	0	0	0
2104	RTA00000527F.i.05.2	37481	2	0	0	0	0	0	0	0
2107	RTA00000523F.k.01.1	41437	1	1	0	0	0	0	0	0
2108	RTA00000425F.j.11.1	76667	1	0	0	0	0	0	0	0
2109	RTA00000424F.b.22.4	72971	1	0	0	0	0	0	0	0
2111	RTA00000525F.a.03.1	36786	2	0	0	0	0	0	0	0
2112	RTA00000527F.i.21.2	37490	2	0	0	0	0	0	0	0
2113	RTA00000424F.a.24.4	73951	1	0	0	0	0	0	0	0
2114	RTA00000522F.k.14.1	74280	1	0	0	0	0	0	0	0
2115	RTA00000522F.n.05.1	73260	1	0	0	0	0	0	0	0
2116	RTA00000523F.c.18.1	66179	1	0	0	0	0	0	0	0

SEQ ID	Sequence Name	cluster	lib 1 clones	lib 2 clones		lib 16 clones			lib 19 clones	lib 20 clones
NO: 2117	RTA00000523F.b.13.1	66330	1	0	0	0	0	0	0	0
2119	RTA00000527F.p.16.1	23798	2	1	0	0	0	0	0	0
2120	RTA00000425F.c.20.1	73581	1	0	0	0	0	0	0	0
2121	RTA00000424F.i.21.1	73482	1	0	0	0	0	0	0	0
2122	RTA00000523F.j.19.1	65910	1	0	0	0	0	0	0	0
2124	RTA00000424F.b.22.1	72971	1	0	0	0	0	0	0	0
2125	RTA00000527F.b.18.1	37469	2	0	0	0	0	0	0	0
2129	RTA00000525F.e.16.1	36837	2	0	0	0	0	0	0	0
2131	RTA00000522F.d.08.1	74284	1	0	0	0	0	0	0	0
2134	RTA00000527F.g.07.1	37488	2	0	0	0	0	0	0	0
2136	RTA00000525F.b.05.1	21116	2	1	0	0	0	0	0	0
2137	RTA00000425F.n.05.1	73965	1	0	0	0	0	0	0	0
2138	RTA00000523F.d.18.1	64072	1	0	0	0	0	0	0	0
2139	RTA00000525F.a.02.1	37454	2	0	0	0	0	0	0	0
2141	RTA00000426F.h.09.1	78797	1	0	0	0	0	0	0	0
2144	RTA00000427F.g.05.1	63138	1	0	0	0	0	0	0	0
2145	RTA00000424F.m.12.1	77675	1	0	0	0	0	0	0	0
2151	RTA00000427F.h.12.1	36894	2	0	0	0	0	0	0	0
2152	RTA00000523F.c.15.1	36935	2	0	0	0	0	0	0	0
2153	RTA00000427F.k.17.1	64965	1	0	0	0	0	0	0	0
2155	RTA00000424F.c.14.3	76614	1	0	0	0	0	0	0	0
2156	RTA00000522F.k.10.2	77619	1	0	0	0	0	0	.0	0
2157	RTA00000424F.m.22.1	72943	1	0	0	0	0	0	0	0
2158	RTA00000527F.h.17.1	37799	2	0	0	0	0	0	0	0
2159	RTA00000527F.c.22.1	37496	2	0	0	0	0	0	0	0
2160	RTA00000425F.k.22.1	78123	1	0	0	0	0	0	0	0
2161	RTA00000424F.m.14.1	77491	i	0	0	0	0	0	0	0
2162	RTA00000522F.k.19.1	32625	1	1	0	0	0	0	0	0
2163	RTA00000523F.i.18.1	64463	1	0	0	0	0	0	0	0
2164	RTA00000425F.j.22.1	73882	1	0	0	0	0	0	0	0
2165	RTA00000527F.g.23.1	37538	2	0	0	0	0	0	0	0
2166	RTA00000426F.m.24.1	63943	1	0	0	0	0	0	0	0
2168	RTA00000425F.d.21.1	78920	1	0	0	0	0	0	0	0
2170	RTA00000424F.d.04.3	76505	1	0	0	0	0	0	0	0
2171	RTA00000424F.d.04.1	76505	1	0	0	0	0	0	0	0
2172	RTA00000427F.c.12.1	66995	1	0	0	0	0	0	0	0
2174	RTA00000527F.I.13.1	36904	2	0	0	0	0	0	0	0
2175	RTA00000522F.h.13.1	40823	1	1	0	0	0	0	0	0
2176	RTA00000424F.l.19.1	75454	1	0	0	0	0	0	0	0
				<b></b>						

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones		lib 19 clones	lib 20 clones
2179	RTA00000427F.a.06.1	66550	1	0	0	0	0	0	0	0
2180	RTA00000525F.c.19.1	38159	2	0	0	0	0	0	0	0
2181	RTA00000523F.f.06.1	62871	1	0	0	0	0	0	0	0
2182	RTA00000424F.h.10.1	72925	1	0	0	0	0	0	0	0
2183	RTA00000522F.a.12.1	33515	1	1	0	0	0	0	0	0
2184	RTA00000522F.h.01.1	75010	1	0	0	0	0	0	0	0
2186	RTA00000425F.e.21.1	77203	1	0	0	0	0	0	0	0
2187	RTA00000523F.f.07.1	62799	1	0	0	0	0	0	0	0
2189	RTA00000424F.j.12.1	73827	1	0	0	0	0	0	0	0
2191	RTA00000523F.d.12.1	64888	1	0	0	0	0	0	0	0
2192	RTA00000523F.e.10.1	62878	1	0	0	0	0	0	0	0
2193	RTA00000425F.f.11.1	79275	1	0	0	0	0	0	0	0
2194	RTA00000426F.m.18.1	62974	1	0	0	0	0	0	0	0
2197	RTA00000522F.g.15.1	76536	1	0	0	0	0	0	0	0
2198	RTA00000522F.n.12.1	74117	1	0	0	0	0	0	0	0
2200	RTA00000424F.d.10.3	73110	1	0	0	0	0	0	0	0
2204	RTA00000527F.c.04.1	23090	3	0	0	0	0	0	0	0
2206	RTA00000527F.h.21.1	37630	2	0	0	0	0	0	0	0
2207	RTA00000425F.c.07.1	76042	1	0	0	0	0	0	0	0
2209	RTA00000525F.c.15.1	7692	2	0	0	0	0	0	0	0
2210	RTA00000424F.d.22.3	76189	1	0	0	0	0	0	0	0
2211	RTA00000523F.h.12.1	65745	1	0	0	0	0	0	- 0	0
2212	RTA00000522F.g.22.1	77504	1	0	0	0	0	0	0	0
2215	RTA00000522F.j.12.2	74341	1	0	0	0	0	0	0	0
2216	RTA00000523F.i.08.1	65099	1	0	0	0	0	0	0	0
2218	RTA00000425F.j.20.1	26760	1	0	0	0	0	0	0	0
2220	RTA00000427F.f.24.1	64572	1	0	0	0	0	0	0	0
2221	RTA00000527F.a.13.1	37740	2	0	0	0	0	0	0	0
2225	RTA00000424F.a.09.4	77833	1	0	0	0	0	0	0	0
2227	RTA00000525F.f.07.1	37500	2	0	0	0	0	0	0	0
2228	RTA00000424F.j.07.1	79211	1	0	0	0	0	0	0	0
2229	RTA00000424F.m.10.1	34251	1	1	0	0	0	0	0	0
2231	RTA00000522F.g.06.1	78221	1	0	0	0	0	0	0	0
2232	RTA00000424F.h.03.1	74447	1	0	0	0	0	0	0	0 ,
2233		74737	1	0	0	0	0	0	0	0
2234	RTA00000427F.c.22.1	63990	1	0	0	0	0	0	0	0
2235		77666	1	0	0	0	0	0	0	0
2236		76982	1	0	0	0	0	0	0	0
2237	RTA00000427F.h.11.1	26494	1	0	0	0	0	0	0	0

SEQ ID	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones	lib 18 clones	lib 19 clones	lib 20 clones
NO: 2238	RTA00000425F.j.16.1	75631	1	0	0	0	0	0	0	0
2240	RTA00000427F.f.17.1	63803	1	0	0	0	0	0	0	0
2241	RTA00000522F.o.18.1	76366	1	0	0	0	0	0	0	0
2242	RTA00000427F.j.22.1	66367	1	0	0	0	0	0	0	0
2242	RTA00000426F.p.10.1	65845	1	0	0	0	0	0	0	0
2243	RTA00000522F.m.02.1	76834	1	0	0	0	0	0	0	0
2247	RTA00000322F.ini.52.1	75921	1	0	0	0	0	0	0	0
2250	RTA00000424F.n.13.1	74942	1	0	0	0	0	0	0	0
	RTA00000424F.g.14.1	74879	1	0	0	0	0	0	0	0
2251 2252	RTA00000424F.g.14.1	64089	1	0	0	0	0	0	0	0
2256	RTA00000420F.g.19.1	64611	1	0	0	0	0	0	0	0
2258	RTA000004277.g.19.11	74938	1	0	0	0	0	0	0	0
	RTA00000522F.c.01.1 RTA00000522F.g.17.1	76486	1	0	0	0	0	0	0	0
2259	RTA00000522F.j.17.1	63610	1	0	0	0	0	0	0	0
2260 2261	RTA00000523F.g.17.1	73410	1	0	0	0	0	0	1	0
2263	RTA00000523F.e.20.1	65164	1	0	0	0	0	0	0	0
		73533	1	0	0	0	0	0	0	0
2264		66665	1	0	0	0	0	0	0	0
2265		75204	1	0	0	0	0	0	0	0
2266 2267		64915	1	0	0	0	0	0	0	0
2268		77373	1	0	0	0	0	0	0	0
2270		41440	_	1	0	0	0	0	. 0	0
2270	RTA000003231.h.21.1	65193	1	0	0	0	0	0	0	0
2271		40841	1	1	0	0	0	0	0	0
2272		76045	_	0	0	0	0	0	0	0
2274		63781	1	0	0	0	0	0	0	0
2275		64799	_	0	0	0	0	0	0	0
2276		66015	_	0	0	0	0	0	0	0
2277		66492		0	0	0	0	0	0	0
2278		75149		0	0	0	0	0	0	0
2279		37820		0	0	0	0	0	0	0
2282	_	63199		0	0	0	0	0	0	0
2284		15939		2	0	0	0	0	0	0
2285		75550		0	0	0	0	0	0	0
2286		63652		0	0	0	0	0	0	0
2287		65613		0	0	0	0	0	0	0
2288		75305		0	0	0	0	0	0	0
2289		62851		0	0	0	0	0	0	0
2291		41446		1	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones			lib 17 clones			
2292	RTA00000527F.l.05.1	13016	4	0	0	1	1	0	0	0
2293	RTA00000426F.m.02.1	66237	1	0	0	0	0	0	0	0
2296	RTA00000522F.1.22.1	75801	1	0	0	0	0	0	0	0
2297	RTA00000427F.h.19.1	63047	1	0	0	0	0	0	0	0
2299	RTA00000522F.g.21.1	77310	1	0	0	0	0	0	0	0
2301	RTA00000522F.g.20.1	77688	1	0	0	0	0	0	0	0
2304	RTA00000425F.k.20.1	74048	1	0	0	0	0	0	0	0
2306	RTA00000522F.b.07.1	78634	1	0	0	0	0	0	0	0
2307	RTA00000426F.g.19.1	63672	1	0	0	0	0	0	0	0
2308	RTA00000525F.d.19.1	36860	2	0	0	0	0	0	0	0
2310	RTA00000427F.d.10.1	40685	1	1	0	0	0	0	0	0
2313	RTA00000424F.a.05.4	77976	1	0	0	0	0	0	0	0
2315	RTA00000424F.a.05.1	77976	1	0	0	0	0	0	0	0
2316	RTA00000522F.I.15.1	74691	1	0	0	0	0	0	0	0
2317	RTA00000425F.e.02.1	76143	1	0	0	0	0	0	0	0
2318	RTA00000525F.c.11.1	37895	2	0	0	0	0	0	0	0
2320	RTA00000522F.c.14.1	75449	1	0	0	0	0	0	0	0
2321	RTA00000424F.m.08.1	19402	1	2	0	0	0	0	0	0
2322	RTA00000527F.f.18.1	37577	2	0	0	0	0	0	0	0
2324	RTA00000522F.a.06.1	73662	1	0	0	0	0	0	0	0
2327	RTA00000522F.d.23.1	73868	1	0	0	0	0	0	0	0
2330	RTA00000523F.j.10.1	63384	1	0	0	0	0	0	0	0
2331	RTA00000527F.p.08.1	36013	2	0	0	0	0	0	0	0
2333	RTA00000426F.f.17.1	66334	1	0	0	0	0	0	0	0
2334	RTA00000523F.j.21.1	36925	2	0	0	0	0	0	0	0
2339	RTA00000523F.a.01.1	74923	1	0	0	0	0	0	0	0
2341	RTA00000427F.j.06.1	63676	1	0	0	0	0	0	0	0
2342	RTA00000424F.m.04.1	79017	1	0	0	0	0	0	0	0
2343	RTA00000523F.i.17.1	65779	1	0	0	0	0	0	0	0
2346	RTA00000525F.c.18.1	24208	2	1	0	0	0	0	0	0
2347	RTA00000527F.e.09.1	37521	2	0	0	0	0	0	0	0
2348	RTA00000424F.j.08.1	73972	1	0	0	0	0	0	0	0
2350	RTA00000527F.c.09.1	64859	1	0	0	0	0	0	0	0
2353	RTA00000523F.c.03.1	36913	2	0	0	0	0	0	0	0
2354	RTA00000427F.k.21.1	62880	1	0	0	0	0	0	0	0
2356	RTA00000427F.d.09.1	66486	1	0	0	0	0	0	0	0
2357	RTA00000426F.n.17.1	66572	1	0	0	0	0	0	0	0
2360	RTA00000426F.m.03.1	66480	1	0	0	0	0	0	0	0
2361	RTA00000424F.h.06.1	77552	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones				
2362	RTA00000425F.d.06.1	77660	1	0	0	0	0	0	0	0
2363	RTA00000427F.e.12.1	62813	1	0	0	0	0	0	0	0
2366	RTA00000426F.n.23.1	18176	1	0	0	0	0	0	0	0
2367	RTA00000522F.m.19.1	41544	1	1	0	0	0	0	0	0
2368	RTA00000522F.a.05.1	32611	1	1	0	0	0	0	0	0
2369	RTA00000427F.i.09.1	65916	1	0	0	0	0	0	0	0
2370	RTA00000424F.j.09.1	74387	1	0	0	0	0	0	0	0
2371	RTA00000424F.n.11.1	73874	1	0	0	0	0	0	0	0
2373	RTA00000527F.e.13.1	37588	2	0	0	0	0	0	0	0
2375	RTA00000425F.j.19.1	77925	1	0	0	0	0	0	0	0
2376	RTA00000522F.g.12.1	78783	1	0	0	0	0	0	0	0
2377	RTA00000523F.a.07.1	75804	1	0	0	0	0	0	0	0
2378	RTA00000425F.e.19.1	73409	1	0	0	0	0	0	0	0
2379	RTA00000425F.n.19.1	78324	1	0	0	0	0	0	0	0
2384	RTA00000427F.k.07.1	63742	1	0	0	0	0	0	0	0
2387	RTA00000522F.a.17.1	79032	1	0	0	0	0	0	0	0
2388	RTA00000527F.I.19.1	36856	2	0	0	0	0	0	0	0
2389	RTA00000424F.i.11.1	41569	1	1	0	0	0	0	0	0
2391	RTA00000424F.d.19.3	73180	1	0	0	0	0	0	0	0
2392	RTA00000522F.j.09.2	78522	1	0	0	0	0	0	0	0
2393	RTA00000424F.m.24.1	77045	1	0	0	0	0	0	0	0
2394	RTA00000522F.j.19.2	76224	1	0	0	0	0	0	- 0	0
2398	RTA00000527F.j.12.2	37503	2	0	0	0	0	0	0	0
2399	RTA00000522F.g.11.1	75432	1	0	0	0	0	0	0	0
2400	RTA00000522F.k.02.2	77622	1	0	0	0	0	0	0	0
2401	RTA00000427F.e.13.1	66080	1	0	0	0	0	0	0	0
2402	RTA00000426F.f.18.1	63271	1	0	0	0	0	0	0	0
2403	RTA00000427F.a.12.1	63377	1	0	0	0	0	0	0	0
2404	RTA00000424F.b.23.4	77322	1	0	0	0	0	0	0	0
2408	RTA00000427F.f.02.1	36822	2	0	0	0	0	0	0	0
2410	RTA00000424F.i.15.1	78043	1	0	0	0	0	0	0	0
2412	RTA00000522F.m.03.1	79194	1	0	0	0	0	0	0	0
2413	RTA00000522F.a.20.1	74070	1	0	0	0	0	0	0	0
2414	RTA00000424F.b.15.4	74958	1	0	0	0	0	0	0	0
2415	RTA00000527F.g.14.1	37532	2	0	0	0	0	0	0	0
2416	RTA00000522F.d.06.1	74809		0	0	0	0	0	0	0
2418		64599		0	0	0	0	0	0	0
2419		22908		0	0	0	0	0	0	0
2421	RTA00000523F.f.17.1	63984	1	0	0	0	0	0	0	0

SEQ ID NO:	Sequence Name	cluster	lib I clones	lib 2 clones		lib 16 clones				lib 20 clones
2423	RTA00000527F.p.24.1	36832	2	0	0	0	0	0	0	0
2424	RTA00000425F.n.17.1	78304	1	0	0	0	0	0	0	0
2426	RTA00000425F.e.07.1	75992	1	0	0	0	0	0	0	0
2428	RTA00000523F.h.08.1	62893	1	0	0	0	0	0	0	0
2429	RTA00000522F.o.10.1	78798	1	0	0	0	0	0	0	0
2430	RTA00000425F.1.10.1	26893	1	0	0	0	0	0	0	0
2431	RTA00000427F.f.16.1	64122	1	0	0	0	0	0	0	0
2434	RTA00000425F.i.10.1	78736	1	0	0	0	0	0	0	0
2435	RTA00000426F.m.12.1	63740	1	0	0	0	0	0	0	0
2436	RTA00000527F.g.12.1	37746	2	0	0	0	0	0	0	0
2439	RTA00000425F.i.18.1	42255	1	1	0	0	0	0	0	0
2441	RTA00000424F.j.13.1	74485	1	0	0	0	0	0	0	0
2445	RTA00000424F.k.10.1	73232	1	0	0	0	0	0	0	0
2446	RTA00000522F.i.07.2	78377	1	0	0	0	0	0	0	0
2448	RTA00000522F.b.08.1	26915	1	0	0	0	0	0	0	0
2449	RTA00000522F.I.08.1	78781	1	0	0	0	0	0	0	0
2450	RTA00000525F.a.14.1	37566	2	0	0	0	0	0	0	0
2451	RTA00000424F.g.08.1	74928	1	0	0	0	0	0	0	0
2452	RTA00000425F.1.09.1	75251	1	0	0	0	0	0	0	0
2453	RTA00000522F.o.20.1	74853	1	0	0	0	0	0	0	0
2454	RTA00000527F.j.04.2	11809	3	1	0	0	0	0	0	0
2456	RTA00000523F.c.13.1	40668	1	1	0	0	0	0	0	0
2457	RTA00000427F.i.21.1	65540	1	0	0	0	0	0	0	0
2459	RTA00000522F.h.02.1	74947	1	0	0	0	0	0	0	0
2460	RTA00000522F.g.10.1	74294	1	0	0	0	0	0	0	0
2464	RTA00000425F.k.16.1	75282	1	0	0	0	0	0	0	0
2465	RTA00000525F.b.09.1	23472	2	1	0	0	0	0	0	0
2466	RTA00000522F.j.08.2	76613	1	0	0	0	0	0	0	0
2468	RTA00000523F.f.19.1	34169	1	1	0	0	0	0	0	0
2469	RTA00000425F.j.18.1	75561	1	0	0	0	0	1	0	0
2470	RTA00000426F.m.04.1	36865	2	0	0	0	0	0	0	0
2471	RTA00000527F.g.21.1	36028	2	0	0	0	0	0	0	0
2473	RTA00000525F.a.22.1	36848	2	0	0	0	0	0	0	0
2474	RTA00000522F.p.22.1	73322	1	0	0	0	0	0	0	0
2475	RTA00000424F.d.12.2	74342	1	0	0	0	0	0	0	0
2476	RTA00000424F.g.24.1	79156	1	0	0	0	0	0	0	0
2477	RTA00000427F.a.10.1	65370	1	0	0	0	0	0	0	0
2478	RTA00000426F.h.20.1	23187	3	0	0	0	0	0	0	0
2479	RTA00000424F.d.12.3	74342	1	0	0	0	0	0	0	0

$(\mathcal{D}_{\mathcal{F}}(x), \mathcal{D}_{\mathcal{F}}(x),  SEQ ID NO:	Sequence Name	cluster	lib 1 clones	lib 2 clones	lib 15 clones	lib 16 clones	lib 17 clones	lib 18 clones	lib 19 clones	lib 20 clones
2480	RTA00000425F.c.03.1	74643	1	0	0	0	0	0	0	0
2481	RTA00000523F.f.16.1	26522	1	0	0	0	0	0	0	0
2482	RTA00000427F.f.15.1	66734	1	0	0	0	0	0	0	0
2485	RTA00000522F.p.18.1	76376	1	0	0	0	0	0	0	0
2493	RTA00000522F.g.18.1	73226	1	0	0	0	0	0	0	0
2495	RTA00000522F.h.05.1	73358	1	0	0	0	0	0	0	0
2497	RTA00000425F.n.16.1	18265	1	0	0	0	0	0	0	0
2498	RTA00000527F.1.21.1	36439	2	0	0	0	0	0	0	0
2501	RTA00000424F.d.17.3	73958	1	0	0	0	0	0	0	0
2502	RTA00000523F.j.02.1	62853	1	0	0	0	0	0	0	0

Table 21. Clones Deposited on January 22, 1999 cDNA ES18 cDNA Library Ref No. cDNA ES17 cDNA ES19 ATCC Accession No. ATCC No. ATCC No. ATCC No. Clone Names in M00001368A:D07 M00001594A:D06 M00003906A:F04 Library M00003917A:D02 M00001613D:H10 M00003908A:F12 M00001596D:E10 M00001673A:A04 M00003914A:G09 M00003868B:G11 M00001592C:G04 M00003915C:H04 M00001599D:A09 M00003917C:D03 M00003905D:B08 M00003791C:E09 M00001619B:A09 M00003908C:G09 M00003870A:C05 M00001593B:E11 M00003914B:A11 M00003916C:C05 M00003922A:D02 M00001605A:E06 M00003861C:H02 M00001608A:D03 M00003959A:A03 M00003931B:A11 M00001616C:A02 M00003905D:C08 M00001679D:B05 M00001617A:D06 M00003908D:D12 M00001679C:D05 M00001595C:E01 M00003901B:H04 M00001616C:A11 M00001687A:G01 M00004031A:E01 M00003945A:E09 M00001608C:E11 M00004029C:C12 M00003908A:H09 M00001610C:E06 M00003911A:F10 M00001649B:G12 M00001612B:D11 M00003914C:F09 M00003813D:H12 M00001618B:E05 M00003963D:B05 M00004087C:D03 M00001621C:C10 M00003986C:E09 M00004269B:C08 M00001647A:H08 M00004031A:F07 M00004348A:A02 M00001631D:B10 M00003907C:C02 M00001679C:D01 M00001608D:E09 M00003911B:F08 M00001490A:E11 M00001641B:C10 M00003914C:H05 M00001387A:E10 M00001641D:E02 M00003918C:C12 M00001397B:G03 M00001630D:H10 M00003914C:C02 M00001441D:E04 M00001585C:D10 M00003914A:E04 M00001352C:G09 M00001560A:H10 M00003903B:D03 M00001370D:A12 M00001573B:C06 M00003905A:F09 M00001387B:A06 M00001660C:D11 M00003867C:E11 M00001397C:A10 M00001641C:C05 M00003870B:B08 M00001536D:G02 M00001578B:B05 M00003879D:A08 M00003895C:A10 M00001587C:C10 M00003891D:B10 M00001464B:B03 M00001590B:C07 M00003901C:A08 M00004370A:G05 M00001554A:E04 M00003903C:C04 M00001490B:H11 M00001570C:G06 M00003905A:F10 M00001530B:D10 M00001576A:B09 M00003906C:D06 M00001579C:E09 M00001582A:H01 M00003907D:A12 M00001587A:H03 M00001582B:E12 M00003905C:G11 M00001457C:H12 M00001615B:F07 M00003914D:D10 M00001535C:E01 M00001571C:A04 M00003972A:G09 M00001561D:C05 M00001573D:D10 M00003975D:C06 M00001589A:C01 M00001576A:F11 M00003905C:B02 M00001664D:G07 M00001579C:G05 M00003907D:F11 M00001565A:H09 M00001582D:A02 M00003914A:G06 M00001381C:B08 M00001589B:E07 M00003914D:E03 M00001395C:F11 M00001575B:B02 M00003972C:F08 M00001429D:F11 M00001578C:G06 M00003976C:D06 M00001449A:F01 M00001591A:B08 M00003907C:C04 M00001391C:H02 M00001607A:F11 M00003905B:C06 M00001429D:H12

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M00001344B:F12

M00001579C:E06

M00001661C:F11

M00001650B:C10

M00004088C:A12

M00004103C:D04

M00004107A:D01

cDNA Library Ref No.		cDNA ES18	cDNA ES19
ATCC Accession No.	ATCC No.	ATCC No.	ATCC No.
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	M00003971A:A06	M00001656B:A08	M00004062A:H06
	M00001346A:E04	M00001662C:B02	M00004075D:C10
	M00001455C:G07	M00001656B:D05	M00004081D:H09
	M00001402D:F02	M00001661C:F10	M00004089A:B08
	M00001438D:C06	M00001663A:C11	M00004103D:F10
	M00001349B:G05	M00001669A:C10	M00004107B:B04
	M00001389C:A08	M00001651B:B12	M00004032C:B02
	M00001439B:A10	M00001653B:E06	M00004078C:F04
	M00001455B:A09	M00001659C:F02	M00004038B:H10
	M00001441B:D11	M00001661B:F03	M00004089A:E02
	M00001453A:B01	M00001663C:F10	M00004096B:F05
	M00001456D:E08	M00001669A:G12	M00004104C:H12
	M00001399A:C03	M00001674D:C10	M00004110D:A10
	M00004496C:H03	M00001651B:E06	M00004036D:F02
	M00004135D:G02	M00001651C:C05	M00004088C:E04
	M00004692A:E07	M00001657C:C07	M00004104D:A04
	M00004374D:E10	M00001662A:C12	M00004107D:E12
	M00004405D:C04	M00001663D:C06	M00004115D:D08
	M00004312B:H07	M00001590B:C05	M00003846A:D03
	M00003976C:A10	M00001483C:G06	M00004072C:F08
	M00004043A:D02	M00001653A:G07	M00004039B:G08
	M00004081C:H06	M00001625B:C10	M00003986D:D02
	M00004050D:A06	M00001626C:D12	M00003914A:B07
	M00001361B:C07	M00001634D:D02	M00003914D:B02
	M00004341B:G03	M00001641C:C06	M00003971B:B07
	M00001342B:E01	M00001642D:F02	M00003978C:A03
	M00004064D:A11	M00001647B:E04	M00003983B:C08
	M00004087A:G08	M00001632B:E05	M00004033D:D07
	M00004344B:H04	M00001639A:C11	M00004072D:H12
	M00004497A:H03	M00001642D:G10	M00004077B:H11
	M00001338C:E10	M00001624A:G11 M00001626C:G08	M00004080A:F01 M00004092C:B03
	M00001366D:E12 M00001390D:E03	M00001626C:G08 M00001672D:D04	M00004092C:B03
	M00001390D:E03 M00001413B:H09	M00001672D:D04 M00001639A:H06	M00004037B:C04 M00004073C:D04
	M00001413B:H09 M00004271B:B06	M00001639A.1100 M00001662C:A04	M00004073C.D04 M00004081A:A08
	M00004271B:B00 M00004151D:E03	M00001602C.A04 M00001641B:B01	M00004081A.A08
	M00004151D:E05	M00001641B.B01 M00001673C:A02	M00004083B.B03
	M00001000B.C04 M00003802D:B11	M00001675C:A02	M00004090C:C07
	M00003802D.B11 M00001579C:E08	M00001659D:D03	M00004080D:B03
	M00001579C.E08	M00001039B:B03	M00004080D:D03
	M00001337D:C08 M00003779B:E12	M00001631D:B03	M00004090C:C10
	M00003779B.E12 M00001638A:D10	M00001671D.E16 M00001652D:A06	M00004102C:D09
	M00001038A:B10	M00001052D:A00 M00001654C:D05	M00004103C:E09
	M00003774A:B03	M000016542:B03	M0000403571:G10 M00003906A:H07
	M00001679A:F01	M00001636A:B07	M00003300A:1107
	M00001679A:101	M00001647B:C09	M00004083B:G03
	M00001653B:E09	M00001033A:C00 M00001482D:A04	M00001073B:E02
	M00001033B.E09 M00001585A:F07	M00001482D:A04 M00001485C:B10	M00003793C:D09
	M00001383A.F07 M00003811D:A12	M00001483C.B10 M00001457D:A07	M00003702B:1109
	M00003811D.A12 M00001653C:F12	M00001457D:A07 M00001461A:E05	M00001678D:C11
	M00001635C.F12	M00001407A:C03	M00001677D:B07
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DATA I II D CAT	DNIA DOLG	-DNIA EC10	-DNA ECIO
cDNA Library Ref No.		cDNA ES18	cDNA ES19
ATCC Accession No.	ATCC No.	ATCC No.	ATCC No.
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	M00003801A:B10	M00001482C:D02	M00001675B:H03
	M00003844C:A08	M00001484D:G05	M00003808D:D04
	M00001636C:C01	M00001459B:D03	M00003752B:C02
	M00001669C:B01	M00001464B:C11	M00003819D:B11
	M00003755A:A09	M00001511A:A05	M00001677D:B02
	M00003798D:H08	M00001477B:C02	M00001694C:G04
	M00001444C:D05	M00001471A:D04	M00003789C:F06
	M00004040B:F10	M00001485C:H10	M00001678C:C06
	M00001355A:C12	M00001485D:E05	M00001675B:D02
	M00001401A:H07	M00001487C:G03	M00003750C:H05
	M00001393B:B09	M00001514A:B04	M00001694A:B12
	M00001409D:F11	M00001530C:G10	M00001677B:H06
	M00001387B:H07	M00001534A:G06	M00001675C:G01
	M00001394C:C11	M00001539A:C12	M00001675B:C01
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	M00001352C:F06	M00001460A:F07	M00001694B:B08
	M00001476D:G03	M00001472C:A01	M00001677B:E06
	M00001399C:D09	M00001481B:A07	M00004037A:E04
	M00001347C:G08	M00001456D:F05	M00003870A:H01
	M00001453D:G12	M00001456D:G11	M00003842C:D11
	M00001382A:F04	M00001477D:F10	M00003828B:F09
	M00001392D:H04	M00001481A:G06	M00003856C:H09
	M00001429C:G12	M00001464A:B03	M00003851A:C10
	M00001454A:C11	M00001469A:G11	M00003841C:E04
	M00001517B:G08	M00001478B:D07	M00003837C:G08
	M00001535A:D02	M00001473A:C11	M00003828B:E07
	M00001352A:E12	M00001457A:G03	M00003772C:B12
	M00001381B:F06	M00001669B:G02	M00001677D:F03
	M00004117A:D11	M00001479D:G06	M00001678B:B12
	M00004217C:D03	M00001473D:B11	M00001678D:G03
	M00004270A:F11	M00001475A:A12	M00001675C:F01
	M00003996A:A06	M00001460A:G07	M00003809A:H04
	M00004056B:D09	M00001464A:D03	M00003771D:G05
	M00004142A:B12	M00001473D:G01	M00001678A:F05
	M00001396D:B03	M00001476D:C05	M00001677B:B06
	M00001370D:E12	M00001484A:A10	M00003794A:E12
	M00001390C:C11	M00001457C:F02	M00003771B:E05
	M00003989A:H11	M00001459B:A12	M00001678A:A11
	M00001426A:A09	M00001464A:E07	M00003805B:C04
	M00004498D:D05	M00001467A:B03	M00001680B:E10
	M00001391B:G12	M00001514A:B08	M00001679B:H07
	M00001391D:D10	M00001464A:B07	M00003904D:B12
	M00001376B:A02	M00001579A:C03	M00003856C:B08
	M00001405B:D07	M00001517A:G08	M00003858D:G06
	M00001368A:A03	M00001530B:G09	M00003870B:F04
	M00001392D:B11	M00001538A:F12	M00003871C:B05
	M00003900D:B10	M00001540C:B03	M00003875A:C04
	M00001494B:C01	M00001547A:F06	M00003901B:A09
	M00001352C:A05	M00001550A:F07	M00003901C:D03
	M00001408B:G06	M00001567B:G11	M00003904C:B06

cDNA Library Ref No. cDNA ES17 ATCC Accession No. ATCC No.

ATCC No. M00004252C:E03 M00003901C:A03 M00004071D:A10 M00001377B:H01 M00003939A:A02 M00004250D:D10 M00004290A:B03 M00003911D:B04 M00004128B:G01 M00004142A:D08 M00003977A:E04 M00004236C:D10 M00004388B:A08 M00004409B:A11 M00003965A:B11 M00003988A:E10 M00004138A:H09 M00003933C:D06 M00004193C:G11 M00004039C:C01 M00003924B:D04 M00004375C:D01

cDNA ES18 ATCC No. M00001572A:A10 M00001575B:G01 M00001487D:C11 M00001577B:A03 M00001539D:E10 M00001587A:F05 M00001560A:F03 M00001569B:G11 M00001573A:A06 M00001575D:A10 M00001583A:D01 M00001587A:F08 M00001590B:B02 M00001553A:E07 M00001560A:H06 M00001589C:A11 M00001538A:C08 M00001531A:H03 M00001548A:G01 M00001531A:H07 M00001542A:E04 M00001487A:F10 M00001503C:G05 M00001511A:G08 M00001539A:H12 M00001542A:F06 M00001549A:F01 M00001514A:A12 M00001516A:D05 M00001546C:C07 M00001549A:H11 M00001538A:D03 M00001544A:C09 M00001546B:F12 M00001550A:D09 M00001487B:F02 M00001513A:G07 M00001530A:F12 M00001538A:D12 M00001587A:G06 M00001551A:D04 M00001485B:C03

cDNA ES19 ATCC No. M00003901C:F09 M00003904D:B10 M00003850D:H11 M00003902B:D06 M00003879A:C01 M00003877D:G05 M00003881D:C12 M00003903A:H09 M00003905A:A06 M00003875D:D09 M00003879B:A06 M00003823D:G05 M00003763A:C01 M00003903B:C02 M00003905A:E07 M00003867A:D12 M00003857C:C09 M00003829C:D10 M00003839D:E02 M00003841C:F03 M00003903D:C06 M00003852D:E08 M00003845D:A09 M00003824A:G10 M00003841C:F06 M00003848A:C09 M00003857C:F11 M00003816C:C01 M00003843A:E08 M00003850A:F06 M00003813B:A11 M00003855C:F10 M00003850D:B05 M00003841D:F06 M00003858B:G05 M00003854D:A12 M00003857C:G01 M00003816C:E09 M00003813A:G04 M00003850D:A05

cDNA Ref No.;	cDNA Ref ES20	cDNA Ref No. ES27	cDNA Ref ES28
ATCC Accession No.	ATCC No.	ATCC No.	ATCC No.
	M00004092D:B11	M00003842B:D09	M00001653B:G07
	M00003981C:F05	M00003845A:H12	M00001654D:G11
	M00004031D:F05	M00003847B:G03	M00001656B:A07
	M00004097B:D03	M00003847C:E09	M00001664B:D06
	M00003986D:G07	M00003853D:G08	M00001664C:H10
	M00004033B:C02	M00003828A:E04	M00001680B:C01
	M00004037B:A04	M00003867C:H09	M00001681A:F03
	M00004092C:B12	M00003822A:F02	M00001684B:G03
	M00005140D:G09	M00003868C:H10	M00001771A:A07
	M00004897D:G05	M00003871A:A05	M00003774C:D02
	M00004960B:D12	M00003879C:G10	M00003754D:D02
	M00005134C:G04	M00003880C:F10	M00001640B:F03
	M00005139A:F01	M00003881D:D06	M00003763B:H01
	M00005176A:C12	M00003884D:G07	M00003812C:A05
	M00005178A:A07	M00003887A:A06	M00003803C:D09
	M00005212A:A02	M00003889A:D10	M00003801B:B10
	M00005229D:H07	M00003889D:B09	M00003798D:E03
	M00004115C:H04	M00003858D:F12	M00003773B:G01
	M00004687A:C03	M00003774B:B08	M00003773B:G01
	M00004900C:E11	M00001680D:D02	M00003771A:G10 M00001452A:E07
	M00004695B:E04	M00001528A:F09	M00001432A.E07
	M00005134D:A06	M00003748A:B07	M00004029B:111 M00003751B:A05
	M00004103B:B07	M00003746A:B07	M00003731B.A03
	M00005177A:B06	M0000105571.700	M00001003B:A11
	M00005178A:A08	M00003761D:E02	M00001579C:B11
	M00004104D:B05	M00003763D:E10	M00001579C:B11
	M00004117B:G01	M00003768A:E02	M00001579C:1110
	M00004900D:B10	M0000370071.E02	M00001579D:G07
	M00005134D:H03	M00003772A:D07	M00001586D:E02
	M00005173C:A02	M0000377211:D07	M00001580D:E02
	M00005177A:H09	M00003778A:D08	M00001587D:A10
	M00005178B:H01	M00003770A:D09	M00001589A.D12
	M00005216C:B09	M00003799A:D09	M00001590C:H08
	M000032166:B09	M00003804A:H04	M00001631B:A11 M00001597A:E12
	M00003626D:B11	M00003804A:1104 M00003806D:G05	M00001397A:E12 M00001649C:B10
	M00005100B:D02	M00003808D:G03	M00001649C:B10
	M00005137A:E01	M00003808C:B03	M00001614A:E06
	M00004119A:A06	M00003817A:E03	M00001613C:D02
	M00004891D:E07	M00003818B:G12	M00001621D:D03
	M00004958B:D01	M00003818B:Q12	M00001623D:G03
	M00005102C:F09	M00003709B:D03	
	M00005102C:107	M00001330A.A09 M00001432A:E06	M00001624C:A06
	M00005174D:H02	M00001432A:E08 M00001381A:D02	M00001630B:A11
	M00005177C:B04	M00001381A:D02	M00001634B:C10
	M00005177C:B04	M00001383A:G04 M00001384C:E03	M00001639D:B07
	M00003218B:D09 M00004102C:F03	M00001384C:E03 M00001384C:F12	M00001573D:F04
	M00004102C.F03 M00004114B:D09	M00001384C:F12 M00001384D:H07	M00001595B:A09
	M00004114B:D09 M00004119D:A07		M00004156B:A12
	M00004119D:A07	M00001385B:F10	M00004319D:G09
	M00004893C:G03	M00001385C:H11	M00004096A:G02
1	WIUUUU4253A:A12	M00001386A:C02	M00004101C:G08

David B. Oak			
cDNA Ref No.;	cDNA Ref ES20	cDNA Ref No. ES27	cDNA Ref ES28
ATCC Accession No.	ATCC No.	ATCC No.	ATCC No.
	M00005134B:E01	M00001372C:F07	M00004102A:H02
	M00004115C:G03	M00001389D:G11	M00004108A:A09
	M00005175B:H04	M00001371D:G01	M00004111D:D11
	M00005214B:D11	M00001392C:D10	M00004115D:C08
	M00004102D:B05	M00001392D:H06	M00004118D:E08
	M00004115A:B12	M00001397B:B09	M00004121C:F06
	M00004119D:H06	M00001398A:G03	M00004131B:H09
	M00004897D:F03	M00001400A:F06	M00004141D:A09
	M00004960B:A09	M00001410B:G05	M00004090A:F09
	M00005134C:E11	M00001413A:F02	M00004146A:C08
	M00005138B:D12	M00001415B:E09	M00004078B:A11
	M00005176A:A05 M00005214C:A09	M00001425A:C11	M00004176B:E08
		M00001386A:D11	M00004188C:A09
	M00004102C:D01	M00001354C:B06	M00004233C:H09
	M00004960B:A08	M00001339D:G02	M00004241D:F11
	M00001476D:A09	M00001660A:C12	M00004246C:A09
	M00001572A:B06 M00005217D:F12	M00001528A:A01	M00004247C:C12
	M00005217D:F12 M00005233A:G08	M00001343D:C04	M00004248B:E08
	M00005236B:F10	M00001347B:E01	M00004257C:H06
	M00005259B:C01	M00001348A:D04	M00004260D:C12
	M00005254D:B08	M00001349C:C05	M00004295B:D02
	M00005254D:B08	M00001350A:D06	M00004040D:F01
	M00003239C:B03	M00001352D:C05	M00004142D:E10
	M00001373A.D00 M00005259D:H08	M00001380C:E05 M00001354B:B10	M00003853D:D03
	M00003239D:1108 M00003813C:D08	M00001334B:B10 M00001380C:F02	M00003860D:H07
	M00003813C:B08	M00001380C:F02 M00001354C:C10	M00003878C:E04
	M00001330B:B00	M00001354C:C10 M00001355B:G11	M00003879A:G05
	M00001596B:C11	M00001355B:G11	M00003880B:C08
	M00004300C:H09	M00001350D:F00	M00003881A:D09 M00003881C:G09
	M00001486D:D12	M00001360D:E11	M0000381C:G09 M00003901B:A05
	M00001585D:F03	M00001361C:H11	M00003901B.A03
	M00001596B:D09	M00001363C:H02	M00003904D.D10 M00003905C:G10
	M00001570D:E06	M00001366D:G02	M00003905E:G10 M00003906B:F12
	M00001582C:E01	M00001369A:H12	M00003900B:112 M00003909A:H04
	M00001586C:E06	M00001352D:D02	M0000390971:1104
	M00001593B:D10	M00001485D:B10	M00003963A:E03
	M00001595C:H11	M00001457B:E03	M00004353C:H07
	M00001596B:H05	M00001457C:C12	M00003919A:A10
	M00001576A:C11	M00001458C:E01	M00003938A:B04
	M00001596C:F09	M00001462B:A10	M00003939C:F04
	M00001567A:H05	M00001464D:F06	M00003946D:C11
	M00001585D:D11	M00001467D:H05	M00003979A:F03
	M00004688A:A02	M00001468B:H06	M00003985C:F01
	M00004927A:E06	M00001505C:H01	M00003997B:G07
	M00005229D:H09	M00001470A:H01	M00003860D:A01
	M00004117B:A12	M00001457A:B07	M00004035A:A04
	M00004187D:G09	M00001479B:A01	M00004042D:H02
	M00005173B:F01	M00001469D:D02	M00004073B:B01
1	M00005218A:G05	M00001487A:A05	M00003946A:H10

D114 D 414			
cDNA Ref No.;	cDNA Ref ES20	cDNA Ref No. ES27	cDNA Ref ES28
ATCC Accession No.	ATCC No.	ATCC No.	ATCC No.
	M00004118A:H08	M00001352C:H02	M00001423D:A09
	M00005134A:D11	M00001488D:C10	M00004314B:G07
	M00005176C:C09	M00001490C:C12	M00001405D:D11
	M00005230D:F06	M00001493B:D09	M00001408A:H04
	M00005234D:B04	M00001504D:D11	M00001408D:D04
	M00005101C:E09	M00001376B:C06	M00001411D:F05
	M00004206A:E02	M00001506B:D09	M00001412A:E04
	M00001570C:A05	M00001511B:C06	M00001413A:F03
	M00005231A:H04	M00001476B:F10	M00001417B:C04
	M00005235A:A03	M00001450D:D04	M00001417D:A04
	M00004118B:B04	M00001433A:G07	M00001418B:F07
	M00005136D:D06	M00001470C:B10	M00001419D:C10
	M00005231C:B01	M00001437D:C04	M00001402B:F12
	M00004153B:B03	M00001447C:C01	M00001423A:G05
	M00004897C:D06	M00001448B:F06	M00001401C:H03
	M00005136D:G06	M00001449D:A06	M00001423D:D12
	M00005212B:A02	M00001433B:H11	M00001424B:H04
	M00005232A:C10	M00001451D:C10	M00001428B:A09
	M00004692A:H10	M00001452A:C07	M00001430A:A02
	M00005101C:B09	M00001453C:A11	M00001432D:F05
	M00004144A:F04	M00001456B:C09	M00001438B:B09
	M00003852B:D11	M00001454B:G03	M00001445B:E04
	M00001660D:E05	M00001454B:G07	M00001445C:A08
	M00003808A:F09	M00001454C:C08	M00001446C:D09
	M00001656A:D10	M00001454C:F02	M00001448A:G09
	M00001671A:H06	M00001454D:D06	M00001449C:H12
	M00003809C:H07	M00001456B:F10	M00001422C:F12
	M00003853C:C06	M00001455D:A09	M00001352C:H10
	M00003860A:A08	M00001455D:A11	M00004375A:H01
	M00003822B:D08	M00001448D:F09	M00004380B:A05
	M00003845A:E12		M00004444B:D11
	M00003854C:C02		M00001338B:E02
	M00003860B:G09		M00001336B:E02
	M00003822B:G01		M00001341A:112
	M00001670A:C11		M00001344A:G07
	M00003852A:B03		M00001345A:G11
	M00003829D:A11		M00001345B:E10
	M00003854C:F01		M00001345C:B01 M00001346B:B07
	M00003856B:C04		M00001340B:B07
	M00003905A:H11		M00001403B:E09
	M00001530A:F11		M00001352B.F04 M00001451C:E01
	M00003840B:E07		M00001451C:E01 M00001361A:H07
	M00003905B:G03		M00001361A:H07
	M00003303B:E08		M00001362B:H06 M00001372C:G12
	M00003846B:E00		M00001372C:G12 M00001375B:G12
	M00003835A:C12		M00001375B:G12 M00001376A:C05
	M00003303B:1103		M00001376B:A08
	M00003820B:B04		M00001376B:A08 M00001377C:E12
	M00003851C.B00		
	M00003833B:C08 M00003829A:F03		M00001382B:F12
	14100003043A.FU3		M00001385A:F12

cDNA Ref No.; ATCC Accession No. ATCC No.

cDNA Ref ES20

M00001638C:G01 M00003845D:B02 M00001653D:G07 M00001578B:A02 M00001590B:H10 M00001595C:A09 M00001596A:E07 M00001607A:B06 M00001607A:D10 M00001652C:B09 M00001671B:F02 M00001632C:D08 M00001638C:H07 M00001652D:B09 M00001614C:E11 M00001633B:B11 M00001651C:A04 M00001639D:G12 M00001671C:F11 M00001638A:B04 M00001637C:H12 M00001669B:H06 M00001639D:F02 M00001590A:C08 M00001636A:C02

M00001614A:A04 M00001639D:G06 cDNA Ref No. ES27 ATCC No.

cDNA Ref ES28 ATCC No. M00001394A:E04 M00001395A:C09 M00001396A:H03 M00001350B:G11

Table 23. Library Deposited on January 22, 1999 cDNA Ref No.;

Table 23. Library Deposited on January 22, 1999			
cDNA Ref No.;	cDNA Library Ref ES29	cDNA Library Ref ES30	
ATCC Accession No.	ATCC No.	ATCC No.	
Clone Names in	M00001449D:B01	M00001594D:B08	
Library	M00001476D:F03	M00001593A:B07	
	M00001456C:B12	M00001594A:C01	
	M00001469B:B01	M00001594A:D08	
	M00001471A:B04	M00001594A:G09	
	M00001472A:D08	M00001595C:B05	
	M00001473A:A07	M00001594B:F12	
	M00001473C:D09	M00001596D:E03	
	M00001475B:C04	M00001594D:C03	
	M00001475C:G11	M00001592C:F11	
	M00001476A:D11	M00001590D:G07	
	M00001476B:D10	M00001595D:A04	
	M00001468A:C05	M00001595D:G03	
	M00001476C:C11	M00001601A:A06	
	M00001467A:H07	M00001590C:F10	
	M00001477B:E02	M00001589B:B08	
	M00001478B:H08	M00001589C:E06	
	M00001479C:E01	M00001611B:A05	
	M00001480A:D03	M00001601A:E02	
	M00001480C:A05	M00001587A:D01	
	M00001481A:H08	M00001591B:B12	
	M00001481B:D09	M00001590B:G08	
	M00001482A:H05	M00001592C:E05	
	M00001482D:H11	M00001591B:B06	
	M00001483C:G09	M00001591D:C07	
	M00001485A:C05	M00001591D:F06	
	M00001476B:F08	M00001592A:E02	
	M00001460A:E11	M00001592A:H05	
	M00001456C:C11	M00001592B:A04	
	M00001457A:C05	M00001587A:B10	
	M00001457A:G12	M00001609D:G10	
	M00001458A:A11	M00005231D:B09	
	M00001458C:D10	M00001614B:E08	
	M00001458D:A01	M00005217C:C01	
	M00001458D:A02	M00001587A:B01	
	M00001458D:C11	M00001613D:B03	
	M00001458D:D01	M00001613A:F03	
	M00001459B:C11	M00001611C:H11	
	M00001468A:H10	M00001611C:C12	
	M00001460A:C10	M00001611B:E06	
	M00001485B:F05	M00001611B:A09	
	M00001460A:H11	M00001610D:D05	
	M00001461A:F05	M00001610B:C07	
	M00001462A:D03	M00001610C:E07	
	M00001464A:B02	M00001610A:E09	
	M00001464A:E10	M00001601A:E12	
	M00001464A:B12	M00001609B:C09	
	M00001405A:D12 M00001465A:C12	M00001608D:D11	
	M00001465A:E10	M00001608B:A09	
	MISSOCITODIA.DIV	1110000100001700	

cDNA Ref No.;	cDNA Library Ref ES29	cDNA Library Ref ES30
ATCC Accession No.	ATCC No.	ATCC No.
	M00001465A:G06	M00001607D:F06
	M00001466A:F08	M00001607B:C05
	M00001467A:C10	M00001606A:H09
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	M00001545A:B12	M00001605A:E09
	M00001535A:D10	M00001605A:A06
	M00001536A:F11	M00001604A:C11
	M00001537A:H05	M00001604A:C07
	M00001539A:E01	M00001604A:B08
	M00001539A:H02	M00001604A:A09
	M00001539B:G07	M00001610A:H05
	M00001539D:B10	M00005214B:A06
	M00001540D:E02	M00005228A:A09
	M00001541B:E05	M00001567A:B09
	M00001542A:G12	M00001561A:D01
	M00001485B:D09	M00001559A:C08
	M00001545A:B10	M00001559A:A11
	M00001533A:G05	M00001558A:G09
	M00001545A:F02	M00001555A:B12
	M00001545A:G05	M00001554A:A08
	M00001546A:D08	M00001552A:H10
	M00001548A:H04	M00001552A:F06
	M00001550A:E07	M00005231C:B07
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	M00001551A:D06	M00001570A:H01
	M00001551A:H06	M00005214D:D10
	M00001551D:H07	M00001570C:G03
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	M00001450A:B08	M00005212D:F08
	M00001544A:F05	M00005212A:D10
	M00001512A:G05	M00005211C:E09
	M00001483B:D04	M00005211A:E09
	M00001485B:H03	M00005210D:C09
	M00001485C:C08	M00005179D:B03
	M00001486B:D07	M00005179B:H02
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	M00001507A:B02	M00001615A:D06
	M00001507A:C05	M00005216B:D02
	M00001507A:E04	M00001579C:A01
	M00001534A:D03	M00001585B:C03
	M00001511A:G01	M00001585B:A06
	M00001533D:A08	M00001584D:H02
	M00001513A:F05	M00001584A:G03
	M00001514A:G03	M00001583D:B08
	M00001516A:D02	M00001583B:F02
	M00001516A:F06	M00001583A:F07
	M00001517A:B11	M00001583A:A05

cDNA Ref No.;

cDNA Library Ref ES29 cDNA Library Ref ES30 ATCC Accession No. ATCC No. ATCC No. M00001529D:C05 M00001582D:F02 M00001530A:A09 M00001582D:B01 M00001530A:E10 M00001582A:A03 M00001532A:C01 M00001579D:H09 M00001532D:A06 M00001567D:B03 M00001485B:D10 M00001579C:H06 M00001511A:A02 M00001585B:F01 M00004249D:B08 M00001579B:F04 M00004185D:E04 M00001579A:E03 M00004188D:G08 M00001578C:F05 M00004197C:F03 M00001577D:H06 M00004198B:D02 M00001577B:F10 M00004204D:C03 M00001576C:G05 M00004208B:F05 M00001575D:D12 M00004208D:B10 M00001575D:B10 M00004210B:B05 M00001575D:A02 M00001362D:H01 M00001573B:G08 M00004216D:D03 M00001573A:E01 M00004167A:H03 M00001572A:B05 M00004275A:B03 M00001571D:F05 M00004285C:A08 M00001579D:F04 M00004316A:G09 M00001636A:F08 M00004465B:D04 M00001643B:E05 M00004493B:D09 M00001642C:G02 M00001347B:H04 M00001642A:F03 M00001351C:B06 M00001641D:C04 M00001360A:G10 M00001641C:H07 M00004216D:C03 M00001641C:F01 M00004076D:D04 M00001641C:D02 M00001484C:A04 M00001641B:F12 M00001456B:G01 M00001634A:B04 M00003972D:C09 M00001636B:G11 M00003974C:E04 M00001649C:D05 M00003979A:E11 M00001636A:C03 M00003983C:F03 M00001635D:D05 M00003989B:F11 M00001635D:C12 M00004031D:B05 M00001635B:H02 M00004177C:A01 M00001635B:H01 M00004076B:G03 M00001634D:G11 M00004167D:A07 M00001634D:D04 M00004078A:A06 M00001634A:H05 M00004085A:B02 M00001641A:A11 M00004107B:A06 M00001638B:E12 M00004111C:E11 M00001640A:H02 M00004130D:H01 M00001614C:E06 M00004157D:B03 M00001636D:F09 M00004159C:F09 M00001637A:A03 M00004162C:A07 M00001637A:A06 M00004135B:G01 M00001637A:E10 M00004040A:G12 M00001637A:F10

cDNA Ref No.;	cDNA Library Ref ES29	cDNA Library Ref ES30
ATCC Accession No.	ATCC No.	ATCC No.
	M00001453B:H12	M00001637C:C06
	M00001448A:E11	M00001644A:H01
	M00001448B:F09	M00001638B:E03
	M00001448B:H05	M00001649A:E11
	M00001448C:E11	M00001638B:F10
	M00001448C:F10	M00001639A:C03
	M00001448D:F12	M00001639A:G07
	M00001449B:B03	M00001639B:H01
	M00001449C:C05	M00001639B:H05
	M00001449D:G10	M00001639C:A09
	M00001448A:B12	M00001639C:C02
	M00001453A:D08	M00001649C:E11
	M00001451B:A04	M00001649C:H10
	M00001454A:F11	M00001637C:E03
	M00001454A:G03	M00001617A:A08
	M00001455A:F04	M00001622A:H12
	M00001455B:E07	M00001621C:H12
	M00001455D:A06	M00001621B:G05
	M00001364B:B06	M00001620D:H02
	M00004117A:G01	M00001620D:G11
	M00001455D:D11	M00001619D:D10
	M00001456B:A06	M00001619C:C07
	M00001451A:C10	M00001619A:E05
	M00001395A:E03	M00001623A:F04
	M00001366D:C06	M00001618A:A03
	M00001365A:H10	M00001618B:D09
	M00001366D:C12	M00001617A:A01
	M00001373D:B03	M00001616D:C11
	M00001453B:F08	M00001615C:G05
	M00001444D:C01	M00001615C:A11
	M00001375B:C06	M00001615B:G07
	M00001392C:D05	M00001633D:H06
	M00001395A:A12	M00001639C:A10
	M00001395A:H02	M00001615B:A09
	M00001397D:G08	M00001615B:G01
	M00001434A:B10	M00001618A;F10
	M00001416A:D09	M00001632C:H07
	M00001433C:F10	M00001633D:D12
	M00001416A:H02	M00001633D:D09
	M00001428D:B10	M00001618A:F08
	M00001428B:D01	M00001633D:G09
	M00001426D:D12	M00001624A:A03
	M00001400C:D02	M00001633C:F09
	M00001427C:D01	M00001633C:H05
	· · · · · · · · · · · · · · · · · · ·	M00001633C:B09
		M00001633A:E06
		M00001633A:E00
		M00001633C:H11
		M00001625D:G10
		M00001631D:G05

cDNA Ref No.;

cDNA Library Ref ES29

ATCC Accession No. ATCC No.

cDNA Library Ref ES30

ATCC No.

M00001629C:E07

M00001629B:B08

M00001626C:E04

M00001626C:C11

M00001632A:B10

M00001624B:B10

M00001633C:A05

M00001625C:G05

Table 24. Clones Deposited on January 22, 1999

cDNA Ref No.; cDNA Ref
ATCC Accession No.
Clone Names in M0000384.
Library M0000384

cDNA Ref ES31 cDNA Ref No. ES32 ATCC No. M00003843A:E04 M00003906A:F12 M00003842C:G03 M00003906B:H06 M00003842A:A03 M00003906C:C05 M00003841D:A04 M00003907A:F01 M00003841B:E06 M00003907B:C03 M00003841C:H11 M00003907B:D05 M00003844A:A11 M00003918A:D08 M00003841C:F01 M00003918A:F09 M00003918C:H10 M00003841C:H08 M00003841C:D07 M00003924A:D08 M00003844D:A07 M00003958B:E11 M00003845D:G08 M00003958B:H08 M00003852C:B06 M00003960A:G07 M00003854B:A07 M00003971B:A10 M00003854B:D04 M00003972D:H02 M00003859D:C05 M00003973C:C03 M00003860B:F11 M00003974B:B11 M00003867B:G07 M00003974D:F02 M00003867B:G08 M00003974D:H04 M00003841B:E03 M00003975C:F07 M00003822D:B10 M00003977C:A06 M00003867D:A06 M00003977C:B03 M00003868B:G06 M00003977D:A03 M00003867B:D10 M00003977D:A06 M00003831C:G05 M00003977D:D04 M00003901C:B01 M00003978D:G04 M00003868C:C07 M00003980A:F04 M00003820A:A08 M00003980B:C11 M00003820B:D07 M00003981C:B04 M00003820B:D10 M00003982A:B12 M00003822D:C06 M00003982C:G04 M00003823B:F07 M00003984D:B08 M00003824C:D07 M00003985B:G04 M00003825B:B10 M00003985D:E10 M00003825B:B11 M00003986B:A08 M00003828A:D05 M00003986C:D09 M00003822D:D04 M00003986D:C08 M00003987B:E12 M00003830C:A03 M00003840D:H10 M00003987B:F08 M00003832A:A09 M00003987C:G03 M00003988D:A08 M00003833B:B03 M00003833B:C12 M00003989C:D03 M00003834B:G04 M00003989C:G05 M00003835A:A09 M00003989D:F12 M00003835B:H11 M00004029B:F01 M00003835D:G06 M00004029C:C05 M00003837C:E05 M00004029C:G10 M00003837C:F10 M00004030D:F11

cDNA Ref ES33 ATCC No. M00005254D:A10 M00005260B:E11 M00005260A:F04 M00005260A:A12 M00005259B:D12 M00005257D:H11 M00005257D:G07 M00005257D:A06 M00005257C:G01 M00005257A:H11 M00005236B:H10 M00005236B:G03 M00005257C:E05 M00001608C:D02 M00001608C:G04 M00001608D:F11 M00001609C:A12 M00001609C:G05 M00001610C:B07 M00001612D:D12 M00001612D:F06 M00001613A:D02 M00001614A:B10 M00001614C:G07 M00001615C:E07 M00001625C:F10 M00001626D:A02 M00001629A:H09 M00001629D:B10 M00001629D:D10 M00001630C:F09 M00001631A:D03 M00001631A:F06 M00001631A:F12 M00001631B:H04 M00001633A:F11 M00001633A:G10 M00001633B:A12 M00001633B:E03 M00001633C:A08 M00001633C:E12 M00001635B:B02 M00001636A:H12 M00001638A:C08 M00001638B:C08 M00001639D:C12 M00001640A:F05

M00001642D:G08

cDNA Ref No.;	cDNA Ref ES31	cDNA Ref No. ES32	cDNA Ref ES33
ATCC Accession No.	ATCC No.	ATCC No.	ATCC No.
	M00003839A:D07	M00004034A:A01	M00001647D:G07
	M00003839D:E11	M00004034C:G02	M00001649A:E10
	M00003829C:H05	M00004034D:E09	M00001650D:D10
	M00003901B:C03	M00004035B:H09	M00001650D:F11
	M00003878C:F06	M00004036D:B04	M00001651C:D11
	M00003878C:G08	M00004036D:B09	M00001651C:G12
	M00003879A:A02	M00004038A:F02	M00001652B:D06
	M00003879A:B08	M00004038D:G06	M00001652D:G02
	M00003879A:C11	M00004039A:C03	M00001652D:G06
	M00003879A:D02	M00004039A:H11	M00001653A:A05
	M00003879B:G02	M00004039B:A05	M00001653D:H07
	M00003880B:D11	M00004039B:E12	M00001654A:E08
	M00003880C:E11	M00004040C:A01	M00001654B:A01
	M00003880C:H03	M00004051D:E01	M00001654C:D10
	M00003901B:F10	M00004072D:F09	M00001654C:G07
	M00003890B:C08	M00004073A:D10	M00001654C:G09
	M00003877C:A11	M00004075B:G09	M00001655C:C07
	M00003819D:B01	M00004076A:D12	M00001655D:E08
	M00003901B:G11	M00004076D:H07	M00001655D:H11
	M00001692A:G06	M00004078A:C11	M00001656A:H12
	M00003903C:C05	M00004078A:E05	M00001656C:C04
	M00003903C:E12	M00004078A:F07	M00001656D:C04
	M00003903D:C12	M00004078B:C11	M00001657C:C11
	M00003903D:D10	M00004078B:F12	M00001657D:A10
	M00003903D:H11	M00004079D:G08	M00001659D:A09
	M00003904A:C04	M00004081A:E02	M00001661D:D05
	M00003904B:C03	M00004081A:G01	M00001664B:E08
	M00003904C:A08	M00004081C:A10	M00001664B:F06
	M00003881B:F10	M00004083A:E08	M00001669B:C12
	M00003871D:G06	M00004083B:C01	M00001669C:B09
	M00003868D:D09	M00004086D:G08	M00001670A:F09
	M00003868D:D11	M00004087B:A12	M00001678C:F09
	M00003870C:A01	M00004087C:A01	M00001693A:H06
	M00003870C:A10	M00004088C:F01	M00003805D:E06
	M00003870C:E10	M00004088D:A11	M00003806C:A06
	M00003871A:A02	M00004088D:B05	M00003809B:A03
	M00003871A:B09	M00004088D:B10	M00003810A:A02
	M00003871A:C11	M00004090B:B04	M00003810B:B11
	M00003871A:G09	M00004090B:H06	M00003810C:B06
	M00003871C:E04	M00004092B:E05	M00003810D:H09
	M00003871C:F12	M00004093C:C02	M00003811C:C02
	M00003878C:D08	M00004096D:H03	M00003813B:F02
	M00003871D:E11	M00004099D:F01	M00003813C:H08
	M00003877C:G12	M00004100B:C07	M00003813D:B12
	M00003875A:A07	M00004103B:E09	M00003813D:C02
	M00003875A:B01	M00004105C:B05	M00003813D:G06
	M00003875B:F12	M00004105C:C08	M00003814B:C01
	M00003875C:A01	M00004107A:A12	M00003817C:A10
	M00003875C:A09	M00004107B:D07	M00003817C:G06
	M00003875C:G02	M00004108B:B02	M00003817D:D12

cDNA Ref No.;	cDNA Ref ES31	cDNA Ref No. ES32	cDNA Ref ES33
ATCC Accession No.	ATCC No.	ATCC No.	ATCC No.
	M00003876B:C05	M00004108D:E07	M00003821A:H09
	M00003876C:D02	M00004108D:G04	M00003822B:G12
	M00003876C:F02	M00004110A:A10	M00003822C:A07
	M00003877B:H10	M00004110B:A07	M00003823C:B01
	M00003868D:B09	M00004118B:A03	M00003823C:C04
	M00003871D:A10	M00004118B:F01	M00003824A:G11
	M00001669D:D06	M00004118D:B05	M00003824B:C09
	M00001661A:B11	M00004119A:C09	M00003824C:A10
	M00001661B:F06	M00004136D:B02	M00003824D:D08
	M00001662A:C07	M00004137A:D06	M00003825B:F10
	M00001662A:G01	M00004139C:A12	M00003825D:F01
	M00001662B:F06	M00004149C:B02	M00003826C:F05
	M00001663C:F12	M00004159C:G12	M00003829A:B08
	M00001664A:F08	M00004169D:B11	M00003829C:E08
	M00001664D:F04	M00004187D:H06	M00003829D:D12
	M00001661A:E06	M00004228C:H03	M00003829D:F03
	M00001669A:B02	M00004244C:G07	M00003830D:B11
	M00001669B:B12	M00004358D:C02	M00003830D:H11
	M00001669C:C08	M00004690A:G08	M00003833D:H08
	M00001675A:G10	M00004891B:D01	M00003833D:H10
	M00001669D:C03	M00004891C:D04	M00003840A:C10
	M00001660B:E03	M00004895B:E12	M00003840B:F05
	M00001669D:F05	M00004895B:G04	M00003840C:C02
	M00001670B:G12	M00004895D:G07	M00003845C:D04
	M00001671A:A10	M00004898C:F03	M00003845D:A04
	M00001671B:G05	M00004899D:G06	M00003845B:7104
	M00001671C:C11	M00004959D:H12	M00003846C:F08
	M00001672D:E08	M00004960A:B08	M00003848B:E07
	M00001673A:G08	M00004960C:E10	M00003848D:G02
	M00001673B:B07	M00005100A:B02	M00003850C:G09
	M00001673B:F07	M00005100A:C01	M00003851A:A06
	M00001673D:D06	M00005101C:E12	M00003851B:D03
	M00001673D:F10	M00005102C:D03	M00003851B:E01
	M00001674A:G07	M00005134B:E08	M00003851C:F09
	M00001692D:B01	M00005139A:H03	M00003851D:H11
	M00001669C:D09	M00005140C:B10	M00003852B:G04
	M00001655C:E01	M00005140D:C06	M00003852C:F07
	M00001649D:A08	M00005178D:H04	M00003853B:C10
	M00001650A:C11	M00005210A:E06	M00003854C:C09
	M00001651A:H11	M00005212B:E01	M00003855A:A01
	M00001652A:A01	M00005212C:C03	M00003855A:F01
	M00001652B:G10	M00005212C:D02	M00003855B:B09
	M00001652D:E05	M00005212C:H02	M00003856A:G04
	M00001652D:E09	M00005212D:D09	M00003856B:A12
	M00001653B:C06	M00005212D:H01	M00003857A:E12
	M00001653B:G10	M00005216A:D09	M00003857A:H10
	M00001653C:D10	M00005216A:H01	M00003857C:E05
	M00001654D:A03	M00005217B:A06	M00003858B:G02
	M00001654D:E12	M00005218A:F09	M00003860D:E06
	M00001654D:E11	M00005228 A . D02	M00003000D:E00

M00005228A:B03

M00003905C:F12

M00001654D:F11

cDNA Ref No.;	cDNA Ref ES31	cDNA Ref No. ES32	cDNA Ref ES33
ATCC Accession No.		ATCC No.	ATCC No.
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	M00001658D:G12	M00005229B:G12	M00003966B:A04
	M00001675C:A04	M00005229B:H04	M00003966C:A12
	M00001660B:D03	M00005229B:H06	M00003966C:F03
	M00001660B:A09	M00005229D:H03	M00003973D:F08
	M00001659D:C09	M00005230B:H09	M00003974D:E01
	M00001659D:B05	M00005232A:H12	M00003974D:H07
	M00001654D:F12	M00005233B:D04	M00003976B:E06
	M00001659A:D12	M00005233D:H07	M00003976B:H07
	M00001655A:B11	M00005235B:F10	M00003978A:E01
	M00001658B:A07	M00005236A:E04	M00003978A:E09
	M00001658A:G09	M00005236A:G10	M00003978C:A12
	M00001657D:A04	M00005236B:A12	M00003980C:E12
	M00001657B:B04	M00001448B:A07	M00003980C:F12
	M00001656B:E01	M00001448B:G07	M00003981A:A07
	M00001660B:E04	M00001448D:E11	M00003981B:B12
	M00001659C:F10	M00001455A:D10	M00003982A:G03
•	M00003808C:A05	M00001455A:E11	M00003982B:C10
	M00001694D:C12	M00001476D:F12	M00003982B:H10
	M00003746C:E02	M00001478A:F12	M00003983A:D02
	M00003779D:E08	M00001482C:F09	M00003983A:F06
	M00003792A:B10	M00001485C:D07	M00003983A:G02
	M00003793D:A11	M00001485C:G06	M00003983D:E08
	M00003794D:G03	M00001485D:A05	M00003983D:H02
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	M00003797A:D06	M00001487C:G09	M00003986C:G11
	M00003797A:G03	M00001530A:B02	M00003986D:H12
	M00003800B:F03	M00001530A:H05	M00004027A:A08
	M00003805A:F02	M00001530D:A11	M00004028A:B10
	M00003806B:C09	M00001539B:B10	M00004028A:G03
	M00001674A:G11	M00001567A:C04	M00004029B:A01
	M00003806D:D11	M00001567A:C11	M00004029B:A06
	M00001693D:E08	M00001567C:B08	M00004029B:G10
	M00003808D:D08	M00001567C:E07	M00004029C:F02
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	M00003809A:F01	M00001570D:E05	M00004030B:A12
	M00003809B:B02	M00001570D:E07	M00004030B:D08
	M00003809B:E10	M00001573B:A06	M00004030C:A08
	M00003813A:B02	M00001573B:H12	M00004030C:C02
	M00003813A:D08	M00001575A:D05	M00004034C:F05
	M00003813B:E09	M00001575B:C01	M00004035B:F05
	M00003814B:C12	M00001576C:H02	M00004036A:A11
	M00003814B:F12	M00001577A:A03	M00004037C:D04
	M00003815C:C06	M00001578B:A06	M00004038A:E05
	M00003815C:D12	M00001579D:F02	M00004038B:D01
,	M00003817B:C04	M00001582C:C04	M00004039C:E02
	M00003806B:G05	M00001582C:G02	M00004039D:B10
	M00001679A:D10	M00001584A:A07	M00004040A:A07
	M00001675C:C03	M00001584D:B06	M00004040A:B04
	M00001675C:D12	M00001584D:C11	M00004040A:C08

cDNA Ref ES33

M00004040B:C05

M00004040B:F07

M00004069A:E12

M00004069C:C08

M00004077A:G12

M00004085B:G01

M00004087A:B05 M00004090D:F12

M00004092C:D08

M00004097C:E03 M00004097C:H08

M00004097D:B05

ATCC No.

cDNA Ref No.; ATCC Accession No.

cDNA Ref ES31 ATCC No. M00001675D:E10 M00001676B:B09 M00001676B:E01 M00001676C:A04 M00001676C:E07 M00001676D:A02 M00001676D:B02 M00001677A:G11 M00001677B:A12 M00001677B:B04 M00001677D:B01 M00001678D:B11 M00001681C:A08 M00003819B:G01 M00001693C:E09 M00001693C:C12 M00001692B:E01 M00001692A:B06 M00001678B:H01 M00001681D:C12 M00001694A:E03 M00001680B:D02 M00001680A:B02 M00001679D:F02 M00001679D:B02 M00001679A:G06

cDNA Ref No. ES32 ATCC No. M00001585D:B12 M00001586C:H07 M00001589D:A01 M00001590D:B04 M00001592B:B02 M00001592D:H02 M00001594C:E05 M00001594C:H03 M00001594D:G11 M00001595A:C07 M00001595A:D12 M00001595A:E07 M00001595B:G07 M00001595B:G10 M00001595B:H11 M00001595C:A01 M00001595C:A05 M00001595C:B12 M00001595C:E05 M00001595C:E09 M00001595D:C11 M00001596A:A02 M00001596A:D01 M00001596C:G05 M00001607A:A01

#### We Claim:

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- 1. A library of polynucleotides, the library comprising the sequence information of at least one of SEQ ID NOS:1-3544, 3546-4510, 4512-4725, 4727-4748, and 4750-5252.
- 2. The library of claim 1, wherein the library is provided on a nucleic acid array.
  - 3. The library of claim 1, wherein the library is provided in a computer-readable format.
- 4. The library of claim 1, wherein the library comprises a differentially expressed polynucleotide comprising a sequence selected from the group consisting of SEQ ID NOS:65, 174, 203, 252, 253, 387, 419, 420, 491, 552, 560, 581, 590, 648, 693, 726, 746, 990, 1095, 1124, 1205, 1354, 1387, 1780, 1899, 1915, 1979, 2007, 2024, 2245, and 2325.
  - 5. The library of claim 1, wherein the library comprises a polynucleotide differentially expressed in a human breast cancer cell, where the polynucleotide comprises a sequence selected from the group consisting of SEQ ID NOS:15, 36, 44, 45, 89, 146, 154, 159, 165, 174, 172, 183, 203, 261, 364, 366, 387, 419, 420, 496, 503, 510, 512, 529, 552, 560, 564, 570, 590, 606, 644, 646, 693, 707, 711, 726, 746, 754, 756, 875, 902, 921, 942, 990, 1095, 1104, 1122, 1131, 1142, 1170, 1184, 1205, 1286, 1289, 1354, 1387, 1435, 1535, 1751, 1764, 1777, 1795, 1860, 1869, 1882, 1890, 1915, 1933, 1934, 1979, 1980, 2007, 2023, 2040, 2059, 2223, 2245, 2300, 2325, 2409, 2462, 2486, 2488, and 2492.
- 6. The library of claim 1, wherein the library comprises a polynucleotide differentially expressed in a human colon cancer cell, where the polynucleotide comprises a sequence selected from the group consisting of SEQ ID NOS:33, 65, 228, 250, 252, 253, 280, 282, 355, 370, 387, 443, 460, 491, 545, 560, 581, 603, 680, 693, 703, 704, 716, 726, 746, 752, 753, 1095, 1104, 1205, 1241, 1264, 1354, 1387, 1401, 1442, 1514, 1734, 1742, 1780, 1851, 1899, 1915, 1954, 2024, 2066, 2262, and 2325.

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- 7. The library of claim 1, wherein the library comprises a polynucleotide differentially expressed in a human lung cancer cell, where the polynucleotide comprises a sequence selected from the group consisting of SEQ ID NOS: 10, 54, 65, 171, 174, 203, 252, 253, 254, 285, 419, 420, 466, 491, 525, 526, 552, 571, 574, 590, 693, 700, 726, 742, 746, 861, 990, 922, 1088, 1288, 1355, 1417, 1422, 1444, 1454, 1570, 1597, 1979, 2007, 2024, 2034, 2038, 2126, and 2245.
- 8. The library of claim 1, wherein the library comprises a polynucleotide differentially expressed in a human cancer cell, where the polynucleotide comprises a sequence selected from the group consisting of SEQ ID NOS:648 and1899.
- 9. An isolated polynucleotide comprising a nucleotide sequence having at least 90% sequence identity to an identifying sequence of SEQ ID NOS:1-3544, 3546-4510, 4512-4725, 4727-4748, and 4750-5252, or a degenerate variant or fragment thereof.

10. The polynucleotide of claim 9, wherein the polynucleotide comprises a sequence of one of SEQ ID NOS:2503, 2504, 2550, 2555, 2578, 2656, 2667, 2712, 2723, 2728, 2738, 2734, 2754, 2758, 2760, 2832, 2835, 2842, 2843, 2849, 2893, 2933, 2956, 2971, 2981, 3009, 3018, 3019, 3046, 3084, 3190, 3129, 3173, 3226, 3227, 3274, 3290, 20 3356, 3365, 3377, 3381, 3390, 3391, 3404, 3407, 3408, 3409, 3418, 3419, 3451, 3597, 3600, 3618, 3632, 3635, 3646, 3648, 3657, 3665, 3669, 3670, 3671, 3656, 3680, 3686, 3695, 3696, 3700, 3710, 3736, 3762, 3763, 3774, 3775, 3791, 3804, 3806, 3836, 3895, 3905, 3919, 3920, 3927, 3936, 3951, 3974, 3998, 4036, 4038, 4044, 4056, 4072, 4117, 4119, 4152, 4153, 4154, 4172, 4175, 4159, 4175, 4205, 4216, 4223, 4228, 4238, 4241, 25 4243, 4251, 4253, 4261, 4263, 4278, 4288, 4322, 4330, 4343, 4359, 4363, 4364, 4365, 4373, 4375, 4384, 4385, 4406, 4409, 4431, 4434, 4441, 4442, 4444, 4455, 4469, 4473, 4477, 4482, 4489, 4495, 4496, 4498, 4525, 4535, 4536, 4540, 4560, 4616, 4562, 4586, 4605, 4629, 4653, 4654, 4658, 4659, 4660, 4661, 4664, 4665, 4668, 4684, 4682, 4688, 4689, 4710, 4718, 4733, 4724, 4733, 4746, 4755, 4760, 4710, 4777, 4785, 4792, 4794, 4801, 4807, 4821, 4822, 4847, 4850, 4854, 4856, 4866, 4885, 4900, 4901, 4905, 4914, 30 4925, 4929, 4931, 4943, 4944, 4959, 5111, 5020, 5041, 5046, 5059, 5083, 5090, 5094, 5102, 5125, 5174, 5197, 5208, 5217, 5237, 5239, 5241, 5243, 5248, and 5252.

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- 11. A recombinant host cell containing the polynucleotide of claim 9.
- 12. An isolated polypeptide encoded by the polynucleotide of claim 9.
- 13. An antibody that specifically binds a polypeptide of claim 12.
  - 14. A vector comprising the polynucleotide of claim 9.
- - 16. A method of detecting differentially expressed genes correlated with a cancerous state of a mammalian cell, the method comprising the step of:

detecting at least one differentially expressed gene product in a test sample derived from a cell suspected of being cancerous, where the gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS:10, 15, 33, 36, 44, 45, 54, 65, 89, 146, 154, 159, 165, 171, 172, 174, 183, 203, 228, 250, 252, 253, 254, 261, 280, 282, 285, 355, 364, 366, 370, 387, 419, 420, 443, 460, 466, 491, 496, 503, 510, 512, 525, 526, 529, 545, 552, 560, 564, 570, 571, 574, 581, 590, 603, 606, 644, 646, 648, 680, 693, 700, 703, 704, 707, 711, 716, 726, 742, 746, 752, 753, 754, 756, 861, 875, 902, 921, 922, 942, 990, 1088, 1095, 1104, 1122, 1131, 1142, 1170, 1184, 1205, 1286, 1288, 1289, 1354, 1355, 1387, 1417, 1435, 1444, 1454, 1535, 1570, 1597, 1734, 1742, 1751, 1764, 1777, 1780, 1795, 1860, 1869, 1882, 1890, 1899, 1915, 1933, 1934, 1954, 1979, 1980, 2007, 2023, 2024, 2034, 2040, 2059, 2126, 2223, 2245, 2262, 2300, 2325, 2409, 2486, 2462, 2488, 2492, 1241, 1264, 1401, 1422, 1442, 1514, 1851, 1915, 2007, 2024, 2038, 2066, and 2245;

wherein detection of the differentially expressed gene product is correlated with a cancerous state of the cell from which the test sample was derived.

17. The method of claim 16, wherein said detecting step is by hybridization of the test sample to a reference array, wherein the reference array comprises an identifying sequence of at least one of SEQ ID NOS: 65, 174, 203, 252, 253, 387, 419, 420, 491, 552,

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560, 581, 590, 648, 693, 726, 746, 990, 1095, 1124, 1205, 1354, 1387, 1780, 1899, 1915, 1979, 2007, 2024, 2325, and 2245.

- 18. The method of claim 16, wherein the cell is a breast tissue derived cell, and the differentially expressed gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS:36, 44, 45, 89, 146, 154, 159, 165, 172, 174, 183, 203, 261, 364, 366, 387, 419, 420, 496, 503, 510, 512, 529, 552, 560,564, 570, 590, 606, 644, 646, 693, 707, 711, 726, 746, 754, 756, 875, 902, 921, 942, 990, 1095, 1104, 1122, 1131, 1142, 1170, 1184, 1205, 1286, 1289, 1354, 1387, 1435, 1535, 1751, 1764, 1777, 1795, 1860, 1869, 1882, 1890, 1915, 1933, 1934, 1979, 1980, 2007, 2023, 2040, 2059, 2223, 2245, 2300, 2325, 2409, 2462, 2486, 2488, and 2492.
  - 19. The method of claim 16, wherein the cell is a colon tissue derived cell, and the differentially expressed gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS:33, 65, 228, 250, 252, 253, 280, 282, 355, 370, 387, 443, 460, 491, 545, 560, 581, 603, 680, 693, 703, 704, 716, 726, 746, 752, 753, 1095, 1104, 1205, 1241, 1264, 1354, 1387, 1401, 1442, 1514, 1734, 1742, 1780, 1851, 1899, 1915, 1954, 2024, 2066, 2262, and 2325.
- 20. The method of claim 16, wherein the cell is a lung tissue derived cell, and the differentially expressed gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS: 10, 54, 65, 171, 174, 203, 252, 253, 254, 285, 419, 420, 466, 491, 525, 526, 552, 571, 574, 590, 693, 700, 726, 742, 746, 861, 922, 990, 1088, 1288, 1355, 1417, 1422, 1444, 1454, 1570, 1597, 1979, 2007, 2024, 2034, 2038, 2126, and 2245.
  - 21. The method of claim 16, wherein the differentially expressed gene product is encoded by a gene corresponding to a sequence of at least one of SEQ ID NOS:648 and 1899.

#### NOVEL HUMAN GENES AND GENE EXPRESSION PRODUCTS II

#### **Abstract**

This invention relates to novel human polynucleotides and variants thereof, their encoded polypeptides and variants thereof, to genes corresponding to these polynucleotides and to proteins expressed by the genes. The invention also relates to diagnostic and therapeutic agents employing such novel human polynucleotides, their corresponding genes or gene products, e.g., these genes and proteins, including probes, antisense constructs, and antibodies.

ATTORNEY'S DOCKET NUMBER 2300-1481CIP

As a below-named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

<u>Nove</u>	I Human Genes and Gene Expression Products II
the sp	pecification of which (check only one item below):
	is attached hereto.
X	was filed as United States application
	Serial No. <u>09/297,648</u>
	on <u>May 4, 1999</u> ,
	and was amended
	on (if applicable).
<u>X</u>	was filed as PCT international application
	Number <u>PCT/US99/01619</u>
	on <u>January 28, 1999</u>
	and was amended under PCT Article 19
	on (if applicable).

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56(a).

I hereby claim foreign priority benefits under Title 35, United States Code, §119 of any foreign application(s) for patent or inventor's certificate or of any PCT international application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign application(s) for patent or inventor's certificate or any PCT international application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed:

DDIOD EODEIGNI/DOT	A DDI IC A TIONICA	AND AND DDIADITY	CLAIMS UNDER 35 U.S.C. 119:
TRIOR TOREIGNICI	AFFLICATIONS	AND ANT FRIORIT	CLAIMS UNDER 33.0.3.C. 119.

COUNTRY (if PCT, indicate "PCT")	APPLICATION NUMBER	DATE OF FILING {day, month, year)	PRIORITY CLAIMED UNDER 35 USC 119
U.S.A.	60/072,910	January 28, 1998	X_YESNO
U.S.A.	60/075,954	February 24, 1998	X_YES _ NO
U.S.A.	60/080,114	March 31, 1998	X_YES _ NO
U.S.A.	60/080,515	April 3, 1998	X_YES _ NO
U.S.A.	60/105,234	October 21, 1998	X_YESNO
U.S.A.	60/105,877	October 27, 1998	X YES _ NO
U.S.A.	60/080,666	April 3, 1998	X_YES _ NO

### Combined Declaration For Patent Application and Power of Attorney (Continued)

(Includes Reference to PCT International Applications)

ATTORNEY'S DOCKET NUMBER 2300-1481CIP

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) or PCT international application(s) designating the United States of America that is/are listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in that/those prior application(s) in the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, §1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this application:

# PRIOR U.S. APPLICATIONS OR PCT INTERNATIONAL APPLICATIONS DESIGNATING THE U.S. FOR BENEFIT UNDER 35 U.S.C. 120:

U.S. APPLICAT	IONS		STATUS (Check one)	
U.S. APPLICATION NUMBER	U.S. FILING DATE	PATENTED	PENDING	ABANDONED
PCT/US99/01619	January 28, 1999		XX	

POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. (List name and registration number)

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I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

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SIGNATURE OF INVENTOR 202

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DATE 9-30-99

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	belief are beli	eved to be true; and furt are punishable by fine or	her that these state imprisonment,	y own knowledge are true and that tements were made with the knowled or both, under section 1001 of Title lity of the application or any patent	dge that verified the	willful false statements and the Eunited States Code, and that
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DATE	10/6/	99	DATE 12/	7/99	DATE	09130/1999
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DATE			DATE 10	125/99	DATE	12/7/99

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212	RESIDENCE & CITIZENSHIP	Moraga CA.		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP U.S.A.
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	to be true; and fine or imprison	further that these statemen	ts were made with n 1001 of Title 18 of	owledge are true and that all statement the knowledge that willful false state of the United States Code, and that su	ments and th	e like so made are punishable by
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DATE	9/24/99		DATE	10/4/99	DATE	12/7/99
SIGNAT	URE OF INVENTOR	₹ 213	SIGNATURE OF IN	VENTOR 214	SIGNATUE	RE OF INVENTOR 215
DATE	· ,·		DATE		DATE	

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218	RESIDENCE & CITIZENSHIP	_{сітү} Sunnyvale		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 678 Durshire Way	<b>y</b>	сіту Sunnyvale		STATE & ZIP CODE/COUNTRY California, 94087
ATT STATE OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE PARTY OF THE P	FULL NAME OF INVENTOR	FAMILY NAME Kita		FIRST GIVEN NAME David		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	Foster City		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 899 Bounty Drive	e, #204,	Foster City		STATE & ZIP CODE/COUNTRY California 94404
	FULL NAME OF INVENTOR	FAMILY NAME Garcia		FIRST GIVEN NAME Veronica		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY Sunnyvale		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP
mental i	POST OFFICE ADDRESS	POST OFFICE ADDRESS 396 Ano Nuevo,	#412	спү Sunnyvale		STATE & ZIP CODE/COUNTRY California 94086
	FULL NAME OF INVENTOR	FAMILY NAME Jones		FIRST GIVEN NAME Lee		SECOND GIVEN NAME William
221	RESIDENCE & CITIZENSHIP	San Jose		STATE OR FOREIGN COUNTRY California	•	COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 396 Ano Nuevo,	#412	CITY Sunnyvale		STATE & ZIP CODE/COUNTRY California 94086
	belief are beli	eved to be true; and furt are punishable by fine or	her that these stat imprisonment, o	y own knowledge are true and that ements were made with the knowledge or both, under section 1001 of Titl ity of the application or any patent	edge that we e 18 of the	rillful false statements and the United States Code, and that
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N. /	, FULL NAME OF INVENTOR	FAMILY NAME Stache-Crain		FIRST GIVEN NAME Birgit		SECOND GIVEN NAME
7	RESIDENCE & CITIZENSHIP	_{СІТҮ} Sunnyvale		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP DE
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 345 South Mary A	Avenue	сіту Sunnyvale		STATE & ZIP CODE/COUNTRY California 94086
	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY		STATE & ZIP CODE/COUNTRY
	FULL NAME OF FAMILY NAME INVENTOR			FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY	• .	COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY		STATE & ZIP CODE/COUNTRY
	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY		STATE & ZIP CODE/COUNTRY
	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY		STATE & ZIP CODE/COUNTRY
	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	СІТУ		STATE OR FOREIGN COUNTRY  CITY		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS				STATE & ZIP CODE/COUNTRY
_	belief are beli	ieved to be true; and furt are punishable by fine or	ther that these star r imprisonment,	y own knowledge are true and that tements were made with the knowled or both, under section 1001 of Title lity of the application or any patent	dge that we are 18 of the	villful false statements and the e United States Code, and that
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	ATURE OF INVENTO	PH .	SIGNATURE OF I	INVENTUR	DATE	NE OF INVENTOR
ATE	RMS\COMDEC.WP	5/96	DATE		PATE	

COMBENED DECLARATION FOR PATENT APPLICATION AND POWER OF ATTORNEY
(Includes Reference to PCT International Applications)

ATTORNEY'S DOCKET NUMBER 2300-1481CIP

As a below-named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

NOVE	Human Genes and Gene Expression Products II
the sp	pecification of which (check only one item below):
	is attached hereto.
X	was filed as United States application
	Serial No. <u>09/297,648</u>
	on <u>May 4, 1999</u>
	and was amended
	on (if applicable).
X	was filed as PCT international application
	Number PCT/US99/01619
	on <u>January 28</u> , 1999
	and was amended under PCT Article 19
	on (if applicable).

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56(a).

I hereby claim foreign priority benefits under Title 35, United States Code, §119 of any foreign application(s) for patent or inventor's certificate or of any PCT international application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign application(s) for patent or inventor's certificate or any PCT international application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed:

PRIOR FOREIGN/PCT APPLICATION(S) AND ANY	PRIORITY CLAIMS UNDER 35 II S.C. 119	
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COUNTRY (if PCT, indicate "PCT")	APPLICATION NUMBER	DATE OF FILING (day, month, year)	PRIORITY CLAIMED UNDER 35 USC 119
U.S.A.	60/072,910	January 28, 1998	X_YESNO
U.S.A.	60/075,954	February 24, 1998	X_YESNO
U.S.A.	60/080,114	March 31, 1998	<u>X_</u> YES NO
U.S.A.	60/080,515	April 3, 1998	X_YES _ NO
U.S.A.	60/105,234	October 21, 1998	X_YESNO
U.S.A.	60/105,877	October 27, 1998	X_YES _ NO
U.S.A.	60/080,666	April 3, 1998	X_YESNO

#### Combined Declaration For Patent Application and Power of Attorney (Continued)

(Includes Reference to PCT International Applications)

ATTORNEY'S DOCKET NUMBER 2300-1481CIP

Direct Telephone Calls to:

(name and telephone number)

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) or PCT international application(s) designating the United States of America that is/are listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in that/those prior application(s) in the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, §1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this application:

### PRIOR U.S. APPLICATIONS OR PCT INTERNATIONAL APPLICATIONS DESIGNATING THE U.S. FOR BENEFIT UNDER 35 U.S.C. 120:

U.S. APPLICAT	ions		STATUS (Check one)	
U.S. APPLICATION NUMBER	U.S. FILING DATE	PATENTED	PENDING	ABANDONED
PCT/US99/01619	January 28, 1999		XX	

POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. (List name and registration number)

Karl Bozicevic, Reg. No. 28,807 Bret E. Field, Reg. No. 37,620 Dianna L. DeVore, Reg. No. 42,484 Carol L. Francis, Reg. No. 36,513 Pamela J. Sherwood, Reg. No. 36,677 Renee A. Fitts, Reg. No. 35,136

Paula A. Borden, Reg. No. 42,344

nd Correspondence to:

Robert P. Blackburn, Reg. No. 30,447 Joseph H. Guth, Reg. No. 31,261 Alisa A. Harbin, Reg. No. 33,895 Charlene A. Launer, Reg. No. 33,035 David P. Lentini, Reg. No. 33,944 Kimberlin L. Morley, Reg. No. 35,391

Chiron ( 4560 H	P. Blackburn Corporation orton Street Ile, California 946	Name: Registration No. Telephone: (650) 327-3400		
Marie 1	FULL NAME OF INVENTOR	FAMILY NAME Williams	FIRST GIVEN NAME Lewis	SECOND GIVEN NAME T.
201	RESIDENCE & CITIZENSHIP	CITY <b>Tiburon</b>	STATE OR FOREIGN COUNTRY California	COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 3 Miroflores	CITY Tiburon	STATE & ZIP CODE/COUNTRY California 94920
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202		CITY <b>Alamo</b>	STATE OR FOREIGN COUNTRY California	COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1470 Lavorna Road	CITY <b>Alamo</b>	STATE & ZIP CODE/COUNTRY California 94507
	FULL NAME OF INVENTOR	FAMILY NAME Innis	FIRST GIVEN NAME Michael	SECOND GIVEN NAME A.
203	RESIDENCE & CITIZENSHIP	CITY Moraga	STATE OR FOREIGN COUNTRY California	COUNTRY OF CITIZENSHIP U.S.A.
		POST OFFICE ADDRESS 315 Constance Place	CITY Moraga	STATE & ZIP CODE/COUNTRY California 94556

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

SIGNATURE OF INVENTOR 201	SIGNATURE OF INVENTOR 202	SIGNATURE OF INVENTOR 203
DATE	DATE	DATE

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,	FULL NAME OF INVENTOR	FAMILY NAME Garcia		FIRST GIVEN NAME Pablo		SECOND GIVEN NAME Dominguez
204	RESIDENCE & CITIZENSHIP	CITY San Francisco		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 882 Chenery Str	eet	стту San Francisco		STATE & ZIP CODE/COUNTRY California 94131
	FULL NAME OF INVENTOR	FAMILY NAME Sudduth-Klinger		FIRST GIVEN NAME Julie		SECOND GIVEN NAME
205	RESIDENCE & CITIZENSHIP	спү Kensington		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP U.S.A
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 280 Lexington Ro	oad	CITY Kensington		STATE & ZIP CODE/COUNTRY California 94707
	FULL NAME OF INVENTOR	FAMILY NAME Reinhard		FIRST GIVEN NAME Christoph		SECOND GIVEN NAME
206	RESIDENCE & CITIZENSHIP	сітү Alameda		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP DE
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1633 Clinton Avenue		стү Alameda		STATE & ZIP CODE/COUNTRY California 94501
	FULL NAME OF INVENTOR	FAMILY NAME Giese		FIRST GIVEN NAME Klaus		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	Berlin DEX		STATE OR FOREIGN COUNTRY  Germany		COUNTRY OF CITIZENSHIP DE
	POST OFFICE ADDRESS	POST OFFICE ADDRESS Chausseetraub 92		Berlin		STATE & ZIP CODE/COUNTRY Germany 10115
	FULL NAME OF INVENTOR	FAMILY NAME Randazzo		FIRST GIVEN NAME FIlippo		SECOND GIVEN NAME
208	RESIDENCE & CITIZENSHIP	спу San Francisco		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 690 Chestnut Street, #403		спү San Francisco		STATE & ZIP CODE/COUNTRY California 94133
	FULL NAME OF INVENTOR	FAMILY NAME Kennedy		FIRST GIVEN NAME Giulia		SECOND GIVEN NAME C.
209	RESIDENCE & CITIZENSHIP	сіту San Francisco		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 360 Castenada Avenue		San Francisco		STATE & ZIP CODE/COUNTRY California 94116
	belief are belie like so made a	eved to be true; and furt re punishable by fine or	her that these state imprisonment, o	own knowledge are true and that ements were made with the knowledge or both, under section 1001 of Title ty of the application or any patent	edge that we are the	rillful false statements and the United States Code, and that
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SIGNATI	URE OF INVENTOR	R 204	SIGNATURE OF IN	VENTOR 205	SIGNATU	RE OF INVENTOR 206
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7	URE OF INVENTO	10	SIGNATURE OF IN	VENTOR 208	SIGNATU	RE OF INVENTOR 209
DATE	Sq. A. 20.	K 1555	DATE		DATE	

ч	FULL NAME OF INVENTOR	FAMILY NAME Pot		FIRST GIVEN NAME David		SECOND GIVEN NAME
210	RESIDENCE & CITIZENSHIP	_{сітү} San Francisco		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP
,	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1565 5th Avenue	e, #102	CITY San Francisco		STATE & ZIP CODE/COUNTRY California 94112
	FULL NAME OF INVENTOR	FAMILY NAME Kassam		FIRST GIVEN NAME Altaf		SECOND GIVEN NAME
211	RESIDENCE & CITIZENSHIP	сітү Oakland		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 2659 Harold Stre	et	сту Oakland		STATE & ZIP CODE/COUNTRY California 94602
	FULL NAME OF INVENTOR	FAMILY NAME Lamson		FIRST GIVEN NAME George		SECOND GIVEN NAME
212	RESIDENCE & CITIZENSHIP	сітү <b>M</b> oraga		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 232 Sandringham Drive		спу <b>Moraga</b>		STATE & ZIP CODE/COUNTRY California 94556
	FULL NAME OF INVENTOR	FAMILY NAME Drmanac		FIRST GIVEN NAME Radoje		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	Palo Alto		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 850 East Greenwich Place		сіту Palo Alto		STATE & ZIP CODE/COUNTRY California 94303
	FULL NAME OF INVENTOR	FAMILY NAME Crkvenjakov		FIRST GIVEN NAME Radomir		SECOND GIVEN NAME
214	RESIDENCE & CITIZENSHIP	CITY Sunnyvale		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP YU
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 762 Haverhill Drive		CITY Sunnyvale		STATE & ZIP CODE/COUNTRY California 94086
	FULL NAME OF INVENTOR	FAMILY NAME Dickson		FIRST GIVEN NAME Mark		SECOND GIVEN NAME
215	RESIDENCE & CITIZENSHIP	спү Hollister		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1411 Gabilan Drive, #B		сіту Hollister		STATE & ZIP CODE/COUNTRY California 95025
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	to be true; and fine or imprisor	further that these stateme	nts were made with on 1001 of Title 18	owledge are true and that all statements the knowledge that willful false stater of the United States Code, and that suc	nents and th	ne like so made are punishable by
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	URE OF INVENTO	R 210	SIGNATURE OF IN	IVENTOR 211	SIGNATU	RE OF INVENTOR 212
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L IZ	FULL NAME OF INVENTOR	FAMILY NAME Drmanac	FIRST GIVEN NAME Snezana	SECOND GIVEN NAME
216	RESIDENCE & CITIZENSHIP	Palo Alto	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 850 East Greenwich Place	CITY Palo Alto	STATE & ZIP CODE/COUNTRY California 94303
	FULL NAME OF INVENTOR	FAMILY NAME Labat	FIRST GIVEN NAME	SECOND GIVEN NAME
217	RESIDENCE & CITIZENSHIP	сіту San Francisco	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1735 Steiner Street	сіту San Francisco	STATE & ZIP CODE/COUNTRY California 94115
	FULL NAME OF INVENTOR	FAMILY NAME Leshkowitz	FIRST GIVEN NAME Dena	SECOND GIVEN NAME
218	RESIDENCE & CITIZENSHIP	Sunnyvale	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP U.S.A.
MERCEN, S. E. Transparen	POST OFFICE ADDRESS	POST OFFICE ADDRESS 678 Durshire Way	сіту Sunnyvale	STATE & ZIP CODE/COUNTRY California, 94087
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219	RESIDENCE & CITIZENSHIP	CITY Foster City	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP U.S.A.
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	FULL NAME OF INVENTOR	FAMILY NAME Garcia	FIRST GIVEN NAME Veronica	SECOND GIVEN NAME
220	RESIDENCE & CITIZENSHIP	Sunnyvale	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 396 Ano Nuevo, #412	city Sunnyvale	STATE & ZIP CODE/COUNTRY California 94086
	FULL NAME OF INVENTOR	FAMILY NAME Jones	FIRST GIVEN NAME Lee	SECOND GIVEN NAME William
221	RESIDENCE & CITIZENSHIP	San Jose	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 396 Ano Nuevo, #412	CITY Sunnyvale	STATE & ZIP CODE/COUNTRY California 94086

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

SIGNATURE OF INVENTOR 216	SIGNATURE OF INVENTOR 217	SIGNATURE OF INVENTOR 218	
DATE	DATE	DATE	
SIGNATURE OF INVENTOR 219	SIGNATURE OF INVENTOR 220	SIGNATURE OF INVENTOR 221	
DATE	DATE	DATE	

بن ر	FULL' NAME OF INVENTOR	FAMILY NAME Stache-Crain		FIRST GIVEN NAME Birgit		SECOND GIVEN NAME
222	RESIDENCE & CITIZENSHIP	CITY Sunnyvale		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP DE
,	POST OFFICE ADDRESS	POST OFFICE ADDRESS 345 South Mary	Avenue	CITY Sunnyvale		STATE & ZIP CODE/COUNTRY California 94086
	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY		STATE & ZIP CODE/COUNTRY
	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
- Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Cardonna de la Card	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY		STATE & ZIP CODE/COUNTRY
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	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS		СІТУ		STATE & ZIP CODE/COUNTRY
# 10 m	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	СІТҮ		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
Adheren Pyr Marie Carlo Marie	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY		STATE & ZIP CODE/COUNTRY
	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY		STATE & ZIP CODE/COUNTRY
	belief are belief like so made a	eved to be true; and furture punishable by fine or	her that these stat imprisonment, o	y own knowledge are true and that ements were made with the knowledge or both, under section 1001 of Tit ity of the application or any patent	edge that w le 18 of the	villful false statements and the United States Code, and that
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SIGNATURE OF INVENTOR 222 SIGNATURE OF II			SIGNATURE OF IN	VENTOR		RE OF INVENTOR
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	TURE OF INVENTO	R	SIGNATURE OF IN	VVENTOR	SIGNATU	RE OF INVENTOR

ATTORNEY'S DOCKET NUMBER 2300-1481CIP

As a below-named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

	Novel Human Genes and Gene Expression Products II					
tne s	e specification of which (check only one item below):					
	is attached hereto.					
X	was filed as United States application					
	Serial No. <u>09/297,648</u>					
	on May 4, 1999					
	and was amended					
	on (if applicable).					
<u>X</u>	was filed as PCT international application					
	Number PCT/US99/01619					
	on <u>January 28, 1999</u>					
	and was amended under PCT Article 19					
	on (if applicable).					

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56(a).

I hereby claim foreign priority benefits under Title 35, United States Code, §119 of any foreign application(s) for patent or inventor's certificate or of any PCT international application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign application(s) for patent or inventor's certificate or any PCT international application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed:

PRIOR FOREIGN/PCT APPLICATION(S) AND ANY PRIORITY	CLAIMS	HNDER 35 H S C 110.	

COUNTRY (if PCT, Indicate "PCT")	APPLICATION NUMBER	DATE OF FILING (day, month, year)	PRIORITY CLAIMED UNDER 35 USC 119
U.S.A.	60/072,910	January 28, 1998	X_YESNO
U.S.A.	60/075,954	February 24, 1998	X_YES _ NO
U.S.A.	60/080,114	March 31, 1998	X_YESNO
U.S.A.	60/080,515	April 3, 1998	X YES _ NO
U.S.A.	60/105,234	October 21, 1998	X YES _ NO
U.S.A.	60/105,877	October 27, 1998	X_YESNO
U.S.A.	60/080,666	April 3, 1998	X_YESNO

### Combined Declaration For Patent Application and Power of Attorney (Continued)

(Includes Reference to PCT International Applications)

ATTORNEY'S DOCKET NUMBER 2300-1481CIP

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) or PCT international application(s) designating the United States of America that is/are listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in that/those prior application(s) in the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, §1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this application:

# PRIOR U.S. APPLICATIONS OR PCT INTERNATIONAL APPLICATIONS DESIGNATING THE U.S. FOR BENEFIT UNDER 35 U.S.C. 120:

U.S. APPLICAT	STATUS (Check one)			
U.S. APPLICATION NUMBER	U.S. FILING DATE	PATENTED	PENDING	ABANDONED
PCT/US99/01619	January 28, 1999		XX	

POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. (List name and registration number)

Karl Bozicevic, Reg. No. 28,807 Bret E. Field, Reg. No. 37,620 Dianna L. DeVore, Reg. No. 42,484 Carol L. Francis, Reg. No. 36,513 Pamela J. Sherwood, Reg. No. 36,677

OZICE 35 Hai alg Alt elepho acsimil	orrespondence to: VIC, FIELD & FRAN milton Avenue, Sui o, California 9430 ne: (650) 327-34 e: (650) 327-323	ite 200 01 00		Direct Telephone Calls to: (name and telephone number)  Name: Registration No. Telephone: (650) 327-3400
	FULL NAME OF INVENTOR	FAMILY NAME Williams	FIRST GIVEN NAME Lewis	SECOND GIVEN NAME T.
	RESIDENCE & CITIZENSHIP	CITY Tiburon	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP U.S.A.
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Annual II	FULL NAME OF INVENTOR	FAMILY NAME Escobedo	FIRST GIVEN NAME Jaime	SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY Alamo	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1470 Lavorna Road	CITY Alamo	STATE & ZIP CODE/COUNTRY California 94507
· · · · · ·	FULL NAME OF INVENTOR	FAMILY NAME Innis	FIRST GIVEN NAME Michael	SECOND GIVEN NAME A.
	RESIDENCE & CITIZENSHIP	CITY Moraga	STATE OR FOREIGN COUNTRY California	COUNTRY OF CITIZENSHIP U.S.A.
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 315 Constance Place	СІТҮ Moraga	STATE & ZIP CODE/COUNTRY California 94556

DATE

SIGNATURE OF INVENTOR 201

like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that

such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

SIGNATURE OF INVENTOR 202

DATE

SIGNATURE OF INVENTOR 203

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	belief are belie like so made a	ved to be true; and furth re punishable by fine or	ner that these state imprisonment, o	own knowledge are true and that ements were made with the knowledge both, under section 1001 of Title ty of the application or any patent	edge that verified that the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the second of the se	willful false statements and the e United States Code, and that
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 360 Castenada Avenue		San Francisco		STATE & ZIP CODE/COUNTRY California 94116
509		San Francisco		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP U.S.A.
		FAMILY NAME Kennedy		FIRST GIVEN NAME Giulia		SECOND GIVEN NAME C.
manus.		POST OFFICE ADDRESS 690 Chestnut Street, #403		CITY San Francisco		STATE & ZIP CODE/COUNTRY California 94133
208 [] [] []		CITY San Francisco		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP U.S.A.
		FAMILY NAME Randazzo		FIRST GIVEN NAME FILLIPPO		SECOND GIVEN NAME
		POST OFFICE ADDRESS Chausseetraub 92		CITY Berlin		STATE & ZIP CODE/COUNTRY Germany 10115
		CITY Berlin		STATE OR FOREIGN COUNTRY Germany		COUNTRY OF CITIZENSHIP DE
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	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1633 Clinton Ave	nue	сіту Alameda		STATE & ZIP CODE/COUNTRY California 94501
206		_{сітү} Alameda		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP DE
		FAMILY NAME Reinhard		FIRST GIVEN NAME Christoph		SECOND GIVEN NAME
		POST OFFICE ADDRESS 280 Lexington Ro	ad	CITY Kensington		STATE & ZIP CODE/COUNTRY California 94707
505		CITY Kensington		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP U.S.A
		FAMILY NAME Sudduth-Klinger		FIRST GIVEN NAME Julie		SECOND GIVEN NAME
		POST OFFICE ADDRESS 882 Chenery Stree		San Francisco		California 94131
Z04		San Francisco		STATE OR FOREIGN COUNTRY  California		CL COUNTRY OF CITIZENSHIP
		FAMILY NAME Garcia		FIRST GIVEN NAME Pablo	<del></del>	SECOND GIVEN NAME Dominguez

	FULL NAME OF INVENTOR	FAMILY NAME Pot	FIRST GIVEN NAME  David	SECOND GIVEN NAME			
210	RESIDENCE & CITIZENSHIP	San Francisco	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP			
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1565 5th Avenue, #102	спу San Francisco	STATE & ZIP CODE/COUNTRY California 94112			
	FULL NAME OF INVENTOR	FAMILY NAME Kassam	FIRST GIVEN NAME Altaf	SECOND GIVEN NAME			
211	RESIDENCE & CITIZENSHIP	спу Oakland	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP U.S.A.			
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 2659 Harold Street	CITY Oakland	STATE & ZIP CODE/COUNTRY  California 94602			
	FULL NAME OF INVENTOR	FAMILY NAME Lamson	FIRST GIVEN NAME George	SECOND GIVEN NAME			
212	RESIDENCE & CITIZENSHIP	спү <b>Moraga</b>	STATE OR FOREIGN COUNTRY  California	COUNTRY OF CITIZENSHIP U.S.A.			
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 232 Sandringham Drive	сіту <b>Mo</b> raga	STATE & ZIP CODE/COUNTRY California 94556			
den den den	FULL NAME OF INVENTOR	FAMILY NAME Drmanac	FIRST GIVEN NAME Radoje	SECOND GIVEN NAME			
213 []] 中 []	RESIDENCE & CITIZENSHIP	Palo Alto	STATE OR FOREIGN COUNTRY California	COUNTRY OF CITIZENSHIP			
# 17 17 17 17 17 17 17 17 17 17 17 17 17	POST OFFICE ADDRESS	POST OFFICE ADDRESS 850 East Greenwich Place	Palo Alto	STATE & ZIP CODE/COUNTRY California 94303			
	FULL NAME OF INVENTOR	FAMILY NAME Crkvenjakov	FIRST GIVEN NAME Radomir	SECOND GIVEN NAME			
214	RESIDENCE & CITIZENSHIP	Sunnyvale CA	STATE OR FOREIGN COUNTRY California	COUNTRY OF CITIZENSHIP			
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 762 Haverhill Drive	CITY Sunnyvale	STATE & ZIP CODE/COUNTRY California 94086			
	FULL NAME OF INVENTOR 15 - CO	FAMILY NAME Dickson	FIRST GIVEN NAME Mark	SECOND GIVEN NAME			
215	RESIDENCE & CITIZENSHIP	Hollister CA.	STATE OR FOREIGN COUNTRY California	COUNTRY OF CITIZENSHIP U.S.A.			
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1411 Gabilan Drive, #B	спу Hollister	STATE & ZIP CODE/COUNTRY California 95025			
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	FULL NAME OF INVENTOR			FIRST GIVEN NAME Birgit		SECOND GIVEN NAME
222	RESIDENCE & CITIZENSHIP	CITY Sunnyvale	<b>A</b>	STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP DE
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 345 South Mary	Avenue	_{СІТҮ} Sunnyvale		STATE & ZIP CODE/COUNTRY  California 94086
	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
-	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
	POST OFFICE ADDRESS	POST OFFICE ADDRESS		СІТУ		STATE & ZIP CODE/COUNTRY
	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
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114 111 21	POST OFFICE ADDRESS	POST OFFICE ADDRESS		CITY		STATE & ZIP CODE/COUNTRY
100	FULL NAME OF INVENTOR	FAMILY NAME		FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP	CITY		STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
egh-controls, Planskings	POST OFFICE ADDRESS	POST OFFICE ADDRESS  FAMILY NAME  CITY		СІТУ		STATE & ZIP CODE/COUNTRY
	FULL NAME OF INVENTOR			FIRST GIVEN NAME		SECOND GIVEN NAME
	RESIDENCE & CITIZENSHIP			STATE OR FOREIGN COUNTRY		COUNTRY OF CITIZENSHIP
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	FULL NAME OF INVENTOR	Drmanac		FIRST GIVEN NAME Snezana		SECOND GIVEN NAME	
216	RESIDENCE & CITIZENSHIP	Palo Alto		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP	
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	FULL NAME OF INVENTOR	FAMILY NAME Labat		FIRST GIVEN NAME Ivan		SECOND GIVEN NAME	
217	RESIDENCE & CITIZENSHIP	San Francisco	<b>-</b> A .	STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP	
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 1735 Steiner Str	eet	сттү San Francisco		STATE & ZIP CODE/COUNTRY California 94115	
	FULL NAME OF INVENTOR	FAMILY NAME Leshkowitz		FIRST GIVEN NAME Dena		SECOND GIVEN NAME	
218	RESIDENCE & CITIZENSHIP	Sunnyvale	<b>A</b> .	STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP U.S.A.	
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 678 Durshire Wa	у	сіту Sunnyvale		STATE & ZIP CODE/COUNTRY California, 94087	
	FULL NAME OF INVENTOR	FAMILY NAME Kita		FIRST GIVEN NAME  David		SECOND GIVEN NAME	
219 ####################################	RESIDENCE & CITIZENSHIP	CITY Foster City		STATE OR FOREIGN COUNTRY California		COUNTRY OF CITIZENSHIP U.S.A.	
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 899 Bounty Drive, #204,		CITY Foster City		STATE & ZIP CODE/COUNTRY California 94404	
Secretaria	FULL NAME OF INVENTOR	FAMILY NAME Garcia		FIRST GIVEN NAME Veronica		SECOND GIVEN NAME	
220	RESIDENCE & CITIZENSHIP	Sunnyvale		STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP ES	
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 396 Ano Nuevo, #412		спү Sunnyvale		state & zip code/country California 94086	
	FULL NAME OF INVENTOR  2 1 - CO	FAMILY NAME Jones		FIRST GIVEN NAME Lee		SECOND GIVEN NAME William	
221	RESIDENCE & CITIZENSHIP	San Jose	<b>\</b>	STATE OR FOREIGN COUNTRY  California		COUNTRY OF CITIZENSHIP U.S.A.	
	POST OFFICE ADDRESS	POST OFFICE ADDRESS 396 Ano Nuevo,	#412	CITY Sunnyvale		STATE & ZIP CODE/COUNTRY California 94086	
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